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(54) Title: VIRAL MATERIAL AND NUCLEOTIDE FRAGMENTS ASSOCIATED WITH MULTIPLE SCLEROSIS, FOR DIAGNOS- TIC, PROPHYLACTIC AND THERAPEUTIC PURPOSES				
(57) Abstract				
<p>The invention relates to a nucleic material, in the isolated or purified state, comprising a nucleotide sequence selected from the group including sequences SEQ ID NO:93, SEQ ID NO:94, their complementary sequences and their equivalent sequences, in particular nucleotide sequences displaying, for any succession of 100 contiguous monomers, at least 50 % and preferably at least 60 % homology with said sequences SEQ ID NO:93, SEQ ID NO:94 and their complementary sequences, excluding HSERV-9 sequence.</p>				

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VIRAL MATERIAL AND NUCLEOTIDE FRAGMENTS ASSOCIATED WITH  
MULTIPLE SCLEROSIS, FOR DIAGNOSTIC, PROPHYLACTIC AND  
THERAPEUTIC PURPOSES

5       Multiple sclerosis (MS) is a demyelinating disease of the central nervous system (CNS) the cause of which remains as yet unknown.

10      "Multiple sclerosis (MS) is the most common neurological disease of young adults with a prevalence in Europe and North America of between 20 and 200 per 100,000. It is characterized clinically by a relapsing/remitting or chronic progressive course, frequently leading to severe disability. Current knowledge suggests that MS is associated with autoimmunity, that 15 genetic background has an important influence and that "infectious" agent(s) may be involved. Indeed, many viruses have been proposed as possible candidates but as yet, none of them has been shown to play an aetiological role.

20      Many studies have supported the hypothesis of a viral aetiology of the disease, but none of the known viruses tested has proved to be the causal agent sought: a review of the viruses sought for several years in MS has been compiled by E. Norrby (1) and R.T. Johnson (2).

25      The discovery of pathogenic retroviruses in man (HTLVs and HIVs) was followed by great interest in their ability to impair the immune system and to provoke central nervous system inflammation and/or degeneration. In the case of HTLV-1, its association with a chronic 30 inflammatory demyelinating disease in man (48) led to extensive investigations to search for an HTLV1-like retrovirus in MS patients. However, despite initial claims, the presence of HTLV-1 or HTLV-like retroviruses was not confirmed.

Recently, a retrovirus different from the known human retroviruses has been isolated in patients suffering from MS (3, 4, and 5).

In 1989, the authors described the production of 5 extracellular virions, associated with reverse transcriptase (RT) activity, by a culture of leptomeningeal cells (LM7) obtained from the cerebrospinal fluid of a patient with MS (3). This was followed by similar findings in monocyte cultures from a series of MS 10 patients (5). Neither viral particles nor viral RT-activity were found in control individuals. Furthermore, the authors were able to transfer the LM7 virus to non-infected leptomeningeal cells *in vitro* (26). The molecular characterization of the "LM7" retrovirus was a 15 prerequisite for further evaluation of its possible role in MS. Considerable difficulties arose from the absence of continuously productive retroviral cultures and from the low levels of expression in the few transient cultures. The strategy described here focused on RNA from 20 extracellular virions, in order to avoid non-specific detection of cellular RNA and of endogenous elements from contaminating human DNA. A specific retroviral sequence associated with virions produced by cell cultures from several MS patients has been identified. The entire 25 sequence of this novel retroviral genome is currently being obtained using RT-PCR on RNA from extracellular virions. The retrovirus previously called "LM7 virus" corresponds to an oncovirus and is now designated MSRV (Multiple Sclerosis-associated RetroVirus).

30 The authors were also able to show that this retrovirus could be transmitted *in vitro*, that patients suffering from MS produced antibodies capable of recognizing proteins associated with the infection of leptomeningeal cells by this retrovirus, and that the 35 expression of the latter could be strongly stimulated by the immediate-early genes of some herpesviruses (6).

All these results point to the role in MS of at least one unknown retrovirus or of a virus having reverse transcriptase activity which is detectable according to the method published by H. Perron (3) and qualified as 5 "LM7-like RT" activity. The content of the publication identified by (3) is incorporated in the present description by reference.

Recently, the Applicant's studies have enabled two continuous cell lines infected with natural isolates 10 originating from two different patients suffering from MS to be obtained by a culture method as described in the document WO-A-93/20188, the content of which is incorporated in the present description by reference. These two lines, derived from human choroid plexus cells, designated 15 LM7PC and PLI-2, were deposited with the ECACC on 22nd July 1992 and 8th January 1993, respectively, under numbers 92072201 and 93010817, in accordance with the provisions of the Budapest Treaty. Moreover, the viral isolates possessing LM7-like RT activity were also 20 deposited with the ECACC under the overall designation of "strains". The "strain" or isolate harboured by the PLI-2 line, designated POL-2, was deposited with the ECACC on 22nd July 1992 under No. V92072202. The "strain" or isolate harboured by the LM7PC line, designated MS7PG, was 25 deposited with the ECACC on 8th January 1993 under No. V93010816.

Starting from the cultures and isolates mentioned above, characterized by biological and morphological criteria, the next step was to endeavour to 30 characterize the nucleic acid material associated with the viral particles produced in these cultures.

The portions of the genome which have already been characterized have been used to develop tests for molecular detection of the viral genome and 35 immunoserological tests, using the amino acid sequences encoded by the nucleotide sequences of the viral genome,

in order to detect the immune response directed against epitopes associated with the infection and/or viral expression.

These tools have already enabled an association 5 to be confirmed between MS and the expression of the sequences identified in the patents cited later. However, the viral system discovered by the Applicant is related to a complex retroviral system. In effect, the sequences to be found encapsidated in the extracellular viral particles 10 produced by the different cultures of cells of patients suffering from MS show clearly that there is coencapsidation of retroviral genomes which are related but different from the "wild-type" retroviral genome which produces the infective viral particles. This phenomenon 15 has been observed between replicative retroviruses and endogenous retroviruses belonging to the same family, or even heterologous retroviruses. The notion of endogenous retroviruses is very important in the context of our discovery since, in the case of MSRV-1, it has been 20 observed that endogenous retroviral sequences comprising sequences homologous to the MSRV-1 genome exist in normal human DNA. The existence of endogenous retroviral elements (ERV) related to MSRV-1 by all or part of their genome explains the fact that the expression of the MSRV-1 25 retrovirus in human cells is able to interact with closely related endogenous sequences. These interactions are to be found in the case of pathogenic and/or infectious endogenous retroviruses (for example some ecotropic strains of the murine leukaemia virus), and in the case of 30 exogenous retroviruses whose nucleotide sequence may be found partially or wholly, in the form of ERVs, in the host animal's genome (e.g. mouse exogenous mammary tumor virus transmitted via the milk). These interactions consist mainly of (i) a trans-activation or coactivation 35 of ERVs by the replicative retrovirus (ii) and "illegitimate" encapsidation of RNAs related to ERVs, or

of ERVs - or even of cellular RNAs - simply possessing compatible encapsidation sequences, in the retroviral particles produced by the expression of the replicative strain, which are sometimes transmissible and sometimes 5 with a pathogenicity of their own, and (iii) more or less substantial recombinations between the coencapsidated genomes, in particular in the phases of reverse transcription, which lead to the formation of hybrid genomes, which are sometimes transmissible and sometimes 10 with a pathogenicity of their own.

Thus, (i) different sequences related to MSRV-1 have been found in the purified viral particles; (ii) molecular analysis of the different regions of the MSRV-1 retroviral genome should be carried out by systematically 15 analyzing the coencapsidated, interfering and/or recombinant sequences which are generated by the infection and/or expression of MSRV-1; furthermore, some clones may have defective sequence portions produced by the retroviral replication and template errors and/or errors 20 of transcription of the reverse transcriptase; (iii) the families of sequences related to the same retroviral genomic region provide the means for an overall diagnostic detection which may be optimized by the identification of invariable regions among the clones expressed, and by the 25 identification of reading frames responsible for the production of antigenic and/or pathogenic polypeptides which may be produced only by a portion, or even by just one, of the clones expressed, and, under these conditions, the systematic analysis of the clones expressed in the 30 region of a given gene enables the frequency of variation and/or of recombination of the MSRV-1 genome in this region to be evaluated and the optimal sequences for the applications, in particular diagnostic applications, to be defined; (iv) the pathology caused by a retrovirus such as 35 MSRV-1 may be a direct effect of its expression and of the proteins or peptides produced as a result thereof, but

also an effect of the activation, the encapsidation or the recombination of related or heterologous genomes and of the proteins or peptides produced as a result thereof; thus, these genomes associated with the expression of  
5 and/or infection by MSRV-1 are an integral part of the potential pathogenicity of this virus, and hence constitute means of diagnostic detection and special therapeutic targets. Similarly, any agent associated with or cofactor of these interactions responsible for the  
10 pathogenesis in question, such as MSRV-2 or the gliotoxic factor which are described in the patent application published under No. FR-2,716,198, may participate in the development of an overall and very effective strategy for the diagnosis, prognosis, therapeutic monitoring and/or  
15 integrated therapy of MS in particular, but also of any other disease associated with the same agents.

In this context, a parallel discovery has been made in another autoimmune disease, rheumatoid arthritis (RA), which has been described in the French Patent  
20 Application filed under No. 95/02960. This discovery shows that, by applying methodological approaches similar to the ones which were used in the Applicant's work on MS, it was possible to identify a retrovirus expressed in RA which shares the sequences described for MSRV-1 in MS, and also  
25 the coexistence of an associated MSRV-2 sequence also described in MS. As regards MSRV-1, the sequences detected in common in MS and RA relate to the pol and gag genes. In the current state of knowledge, it is possible to associate the gag and pol sequences described with the  
30 MSRV-1 strains expressed in these two diseases.

The present patent application relates to various results which are additional to those already protected by the following French Patent Applications:

- No. 92/04322 of 03.04.1992, published under  
35 No. 2,689,519;

- No. 92/13447 of 03.11.1992, published under  
No. 2,689,521;  
- No. 92/13443 of 03.11.1992, published under  
No. 2,689,520;  
5 - No. 94/01529 of 04.02.1994, published under  
No. 2,715,936;  
- No. 94/01531 of 04.02.1994, published under  
No. 2,715,939;  
- No. 94/01530 of 04.02.1994, published under  
10 No. 2,715,936;  
- No. 94/01532 of 04.02.1994, published under  
No. 2,715,937;  
- No. 94/14322 of 24.11.1994, published under  
No. 2,727,428;  
15 - and No. 94/15810 of 23.12.1994; published under  
No. 2,728,585.

The present invention relates, in the first place, to a viral material, in the isolated or purified state, which may be recognized or characterized in  
20 different ways:

- its genome comprises a nucleotide sequence chosen from the group including the sequences SEQ ID NO:46, SEQ ID NO:51, SEQ ID NO:52, SEQ ID NO:53, SEQ ID NO:56, SEQ ID NO:58, SEQ ID NO:59, SEQ ID NO:60, SEQ ID NO:61, SEQ ID  
25 NO:89, their complementary sequences and their equivalent sequences, in particular nucleotide sequences displaying, for any succession of 100 contiguous monomers, at least 50% and preferably at least 70% homology with the said sequences SEQ ID NO:46, SEQ ID NO:51, SEQ ID NO:52, SEQ ID  
30 NO:53, SEQ ID NO:56, SEQ ID NO:58, SEQ ID NO:59, SEQ ID NO:60 SEQ ID NO:61, SEQ ID NO:89, respectively, and their complementary sequences;

- the region of its genome comprising the env and pol genes and a portion of the gag gene, excluding the  
35 subregion having a sequence identical or equivalent to SEQ ID NO:1, codes for any polypeptide displaying, for any

contiguous succession of at least 30 amino acids, at least 50% and preferably at least 70% homology with a peptide sequence encoded by any nucleotide sequence chosen from the group including SEQ ID NO:46, SEQ ID NO:51, SEQ ID 5 NO:52, SEQ ID NO:53, SEQ ID NO:56, SEQ ID NO:58, SEQ ID NO:59, SEQ ID NO:60 SEQ ID NO:61 SEQ ID NO:89 and their complementary sequences;

- the pol gene comprises a nucleotide sequence partially or totally identical or equivalent to SEQ ID NO:57 or SEQ 10 ID NO:93, excluding SEQ ID NO:1.

- the gag gene comprises a nucleotide sequence partially or totally identical or equivalent to SEQ ID NO:88.

As indicated above, according to the present invention, the viral material as defined above is 15 associated with MS. And as defined by reference to the pol or gag gene of MSRV-1, and more especially to the sequences SEQ ID NOS 51, 56, 57, 59, 60, 61, 88, 89, 93, 169, 170, 171, 172, 176, 177, 178 and 179, this viral material is associated with RA.

20 The present invention also relates to a nucleic material, in the isolated or purified state, having at least one of the following definitions :

- a nucleic material comprising a nucleotide sequence selected from the group including sequences SEQ ID NO:93, 25 SEQ ID NO:94, SEQ ID NO:169, SEQ ID NO:170, SEQ ID NO:171, SEQ ID NO:172, SEQ ID NO:176, SEQ ID NO:177, SEQ ID NO:178, SEQ ID NO:179, their complementary sequences and their equivalent sequences, in particular nucleotide sequences displaying, for any succession of 100 30 contiguous monomers, at least 50% and preferably at least 60% homology with said sequences SEQ ID NO:93, SEQ ID NO:94, SEQ ID NO:169, SEQ ID NO:170, SEQ ID NO:171, SEQ ID NO:172, SEQ ID NO:176, SEQ ID NO:177, SEQ ID NO:178, SEQ ID NO:179, and their complementary 35 sequences, excluding HSERV-9 (or ERV-9) ; advantageously, the nucleotide sequence of said nucleic material is

selected from the group including sequences SEQ ID NO:93, SEQ ID NO:94, SEQ ID NO:169, SEQ ID NO:170, SEQ ID NO:171, SEQ ID NO:172, SEQ ID NO:176, SEQ ID NO:177, SEQ ID NO:178, SEQ ID NO:179, their complementary 5 sequences and their equivalent sequences, in particular nucleotide sequences displaying, for any succession of 100 contiguous monomers, at least 70% and preferably at least 80% homology with said sequences SEQ ID NO:93, SEQ ID NO:94, SEQ ID NO:169, SEQ ID NO:170, SEQ ID NO:171, 10 SEQ ID NO:172, SEQ ID NO:176, SEQ ID NO:177, SEQ ID NO:178, SEQ ID NO:179, and their complementary sequences ;

- a nucleic material, in the isolated or purified state, coding for any polypeptide displaying, for any contiguous 15 succession of at least 30 amino acids, at least 50%, preferably at least 60 %, and most preferably at least 70% homology with a peptide sequence encoded by any nucleotide sequence selected from the group including SEQ ID NO:93, SEQ ID NO:94, SEQ ID NO:169, SEQ ID NO:170, SEQ ID NO:171, 20 SEQ ID NO:172, SEQ ID NO:176, SEQ ID NO:177, SEQ ID NO:178, SEQ ID NO:179 and their complementary sequences;

- a nucleic material, in the isolated or purified state, of retroviral type, comprising a nucleotide sequence 25 identical or similar to at least part of the pol gene of an isolated retrovirus associated with multiple sclerosis or rheumatoid arthritis; advantageously, said nucleotide sequence is 80 % similar to said at least part of the gene pol;

30 - a nucleic material comprising a nucleotide sequence identical or similar to at least part of the pol gen of an isolated virus encoding a reverse transcriptase having a enzymatic site comprised between the amino acid domains LPQG-YXDD, having a phylogenetic distance with HSERV-9 of 35 0.063 ± 0.1, and preferably 0.063 ± 0.05; the phylogenetic distances are calculated on the basis of a reference

sequence according to UPGM tree option of the Geneworks™ Software (INTELLIGENETICS) ;

By enzymatic site, we understand the amino acids domain(s) conferring the specific activity of a given enzyme.

5 The present invention also relates to different nucleotide fragments each comprising a nucleotide sequence chosen from the group including:

- (a) all the genomic sequences, partial and total, of the pol gene of the MSRV-1 virus, except for the total 10 sequence of the nucleotide fragment defined by SEQ ID NO:1;
- (b) all the genomic sequences, partial and total, of the env gene of MSRV-1;
- (c) all the partial genomic sequences of the gag gene of 15 MSRV-1;
- (d) all the genomic sequences overlapping the pol gene and the env gene of the MSRV-1 virus, and overlapping the pol gene and the gag gene;
- (e) all the sequences, partial and total, of a clone 20 chosen from the group including the clones FBd3 (SEQ ID NO:46), t pol (SEQ ID NO:51), JLBC1 (SEQ ID NO:52), JLBC2 (SEQ ID NO:53) and GM3 (SEQ ID NO:56), FBd13 (SEQ ID NO:58), LB19 (SEQ ID NO:59), LTRGAG12 (SEQ ID NO:60), FP6 (SEQ ID NO:61), G+E+A 25 (SEQ ID NO:89), excluding any nucleotide sequence identical to or lying within the sequence defined by SEQ ID NO:1;
- (f) sequences complementary to the said genomic sequences;
- (g) sequences equivalent to the said sequences (a) to (e), 30 in particular nucleotide sequences displaying, for any succession of 100 contiguous monomers, at least 50% and preferably at least 70% homology with the said sequences (a) to (d), provided that this nucleotide fragment does not comprise 35 or consist of the sequence ERV-9 as described in LA MANTIA et al. (18).

The term genomic sequences, partial or total, includes all sequences associated by coencapsidation or by coexpression, or recombined sequences.

Preferably, such a fragment comprises:

- 5 - either a nucleotide sequence identical to a partial or total genomic sequence of the pol gene of the MSRV-1 virus, except for the total sequence of the nucleotide fragment defined by SEQ ID NO:1, or identical to any sequence equivalent to the said partial or total genomic  
10 sequence, in particular one which is homologous to the latter;
- or a nucleotide sequence identical to a partial or total genomic sequence of the env gene of the MSRV-1 virus, or identical to any sequence complementary to the said  
15 nucleotide sequence, or identical to any sequence equivalent to the said nucleotide sequence, in particular one which is homologous to the latter.

In particular, the invention relates to a nucleotide fragment comprising a coding nucleotide sequence which is partially or totally identical to a nucleotide sequence chosen from the group including:

- 20 - the nucleotide sequence defined by SEQ ID NO:40, SEQ ID NO:62 or SEQ ID NO:89;
- sequences complementary to SEQ ID NO:40, SEQ ID NO:62 or  
25 SEQ ID NO:89;
- sequences equivalent, and in particular homologous to SEQ ID NO:40, SEQ ID NO:62 or SEQ ID NO:89;
- sequences coding for all or part of the peptide sequence defined by SEQ ID NO:39, SEQ ID NO:63 or SEQ ID NO:90;
- 30 - sequences coding for all or part of a peptide sequence equivalent, in particular homologous to SEQ ID NO:39, SEQ ID NO:63 or SEQ ID NO:90, which is capable of being recognized by sera of patients infected with the MSRV-1 virus, or in whom the MSRV-1 virus has been reactivated.

The invention also relates to a nucleotide fragment (called fragment I) having at least one of the following definitions :

- a nucleotide fragment comprising a nucleotide sequence  
5 selected from the group including SEQ ID NO:93, SEQ ID NO:94, SEQ ID NO:169, SEQ ID NO:170, SEQ ID NO:171, SEQ ID NO:172, SEQ ID NO:176, SEQ ID NO:177, SEQ ID NO:178, SEQ ID NO:179, their complementary sequences and their equivalent sequences, in particular  
10 nucleotide sequences displaying, for any succession of 100 contiguous monomers, at least 50% and preferably at least 60% homology with said sequences and their complementary sequences, said group excluding SEQ ID NO:1, said nucleotide fragment not comprising nor consisting of  
15 the sequence HSERV-9 (or ERV-9); preferably the nucleotide sequence of said fragment is selected from the group including SEQ ID NO:93, SEQ ID NO:94, SEQ ID NO:169, SEQ ID NO:170, SEQ ID NO:171, SEQ ID NO:172, SEQ ID NO:176, SEQ ID NO:177, SEQ ID NO:178,  
20 SEQ ID NO:179, their complementary sequences and their equivalent sequences, in particular nucleotide sequences displaying, for any succession of 100 contiguous monomers, at least 70% and preferably at least 80% homology with said sequences and their complementary sequences;
- 25 - a nucleotide fragment comprising a coding nucleotide sequence which is partially or totally identical to a nucleotide sequence selected from the group including :
  - SEQ ID NO:93, SEQ ID NO:94, SEQ ID NO:169,
  - SEQ ID NO:170, SEQ ID NO:171, SEQ ID NO:172,
  - 30 SEQ ID NO:176, SEQ ID NO:177, SEQ ID NO:178, SEQ ID NO:179 ; their complementary sequences ; their equivalent sequences, in particular homologous to SEQ ID NO:93, SEQ ID NO:94, SEQ ID NO:169, SEQ ID NO:170, SEQ ID NO:171, SEQ ID NO:172, SEQ ID NO:176,
  - 35 SEQ ID NO:177, SEQ ID NO:178, SEQ ID NO:179;

sequences encoding all or parts of the peptide sequence defined by SEQ ID NO:95, SEQ ID NO:173, SEQ ID NO:174, SEQ ID NO:175, SEQ ID NO:180, SEQ ID NO:181, SEQ ID NO:182;

5 sequences encoding all or parts of a peptide sequence equivalent, in particular homologous to SEQ ID NO:95, SEQ ID NO:173, SEQ ID NO:174, SEQ ID NO:175, SEQ ID NO:180, SEQ ID NO:181, SEQ ID NO:182, which is capable of being recognized by sera of patients infected  
10 with the MSRV-1 virus, or in whom the MSRV-1 virus has been reactivated.

The invention also relates to any nucleic acid probe for the detection of virus associated with MS and/or rheumatoid arthritis (RA), which is capable of hybridizing...  
15 specifically with any fragment such as is defined above, belonging or lying within the genome of the said pathogenic agent. It relates, in addition, to any nucleic acid probe for detection of a pathogenic and/or infective agent associated with RA, which is capable of hybridizing  
20 specifically with any fragment as defined above by reference to the pol and gag genes, and especially with respect to the sequences SEQ ID NOS 40, 51, 56, 59, 60, 61, 62, 89 and SEQ ID NOS 39, 63 and 90.

The invention also relates to a primer for the  
25 amplification by polymerization of an RNA or a DNA of a viral material, associated with MS and/or RA, comprising a nucleotide sequence identical or equivalent to at least one portion of the nucleotide sequence of any fragment such as is defined above, in particular a nucleotide sequence displaying, for any succession of at least 10 contiguous monomers, preferably 15 contiguous monomers, more preferably 18 contiguous monomers and even most preferably 20 contiguous monomers, at least 70% homology with at least the said portion of the said fragment.  
30 Preferably, the nucleotide sequence of such a primer is identical to any one of the sequences selected from the

group including SEQ ID NO:47 to SEQ ID NO:50, SEQ ID NO:55, SEQ ID NO:64, SEQ ID NO:86, SEQ ID NO:99 to SEQ ID NO:111, SEQ ID NO:183, SEQ ID NO:184, SEQ ID NO:185, SEQ ID NO:186.

5 Generally speaking the invention also encompasses any RNA or DNA, and in particular replication vector, comprising a genomic fragment of the viral material such as is defined above, or a nucleotide fragment such as is defined above.

10 The invention also relates to the different peptides encoded by any open reading frame belonging to a nucleotide fragment such as is defined above, in particular any polypeptide, for example any oligopeptide forming or comprising an antigenic determinant recognized 15 by sera of patients infected with the MSRV-1 virus and/or in whom the MSRV-1 virus has been reactivated. Preferably, this polypeptide is antigenic, and is encoded by the open reading frame beginning, in the 5'-3' direction, at nucleotide 181 and ending at nucleotide 330 of 20 SEQ ID NO:1.

The invention also encompasses the following polypeptides :

a)

- a polypeptide encoded by any open reading frame 25 belonging to a nucleotide fragment, fragment I, as defined above ;  
- a polypeptide, characterized in that the open reading frame encoding it, is comprised, in the 5'-3' direction, between nucleotide 18 and nucleotide 2304 of SEQ ID NO:93;  
30 - a polypeptide, having a peptide sequence comprising a sequence partially or totally identical to SEQ ID NO:95;

b)

- a polypeptide, recombinant or synthetic, having a peptide sequence which comprises a sequence identical or 35 equivalent to SEQ ID NO:96; in particular said polypeptide

exhibits an enzymatic activity consisting of proteolytic activity;

- a polypeptide, recombinant or synthetic, characterized in that the open reading frame encoding it begins, in the 5' 5'-3' direction, at nucleotide 18 and ends at nucleotide 340 of SEQ ID NO:93;

- a polypeptide having an inhibitory activity on the proteolytic activity of a polypeptide as defined according to b);

10 c)

- a polypeptide, recombinant or synthetic, having a peptide sequence which comprises a sequence identical or equivalent to SEQ ID NO:97; in particular said polypeptide exhibits a reverse transcriptase activity;

15 - a polypeptide having a peptide sequence which comprises a sequence identical or equivalent to SEQ ID NO:98; in particular said polypeptide exhibits a ribonuclease activity;

20 - a polypeptide, recombinant or synthetic, characterized in that the open reading frame encoding it begins, in the 5' 5'-3' direction, at nucleotide 341 and ends at nucleotide 2304 of SEQ ID NO:93;

- a polypeptide, recombinant or synthetic, characterized in that the open reading frame encoding it begins, in the 25 5' 5'-3' direction, at nucleotide 1858 and ends at nucleotide 2304 of SEQ ID NO:93.

- a polypeptide having an inhibitory activity on the reverse transcriptase activity of a polypeptide as defined according to c) or on the ribonuclease H activity of a 30 polypeptide as defined according to c).

In particular, the invention relates to an antigenic polypeptide recognized by the sera of patients infected with the MSRV-1 virus, and/or in whom the MSRV-1 virus has been reactivated, whose peptide sequence is 35 partially or totally identical or is equivalent to the sequence defined by SEQ ID NO:39, SEQ ID NO:63,

SEQ ID NO:87, SEQ ID NO:95, SEQ ID NO:96, SEQ ID NO:97, SEQ ID NO:98, SEQ ID NO:173, SEQ ID NO:174, SEQ ID NO:175, SEQ ID NO:180, SEQ ID NO:181 and SEQ ID NO:182; such a sequence is identical, for example, to any sequence 5 selected from the group including the sequences SEQ ID NO:41 to SEQ ID NO:44, SEQ ID NO:63 and SEQ ID NO:87.

The present invention also proposes mono- or polyclonal antibodies directed against the MSRV-1 virus, 10 which are obtained by the immunological reaction of a human or animal body or cells to an immunogenic agent consisting of an antigenic polypeptide such as is defined above.

The invention next relates to:

15 - reagents for detection of the MSRV- virus, or of an exposure to the latter, comprising, at least one reactive substance selected from the group consisting of a probe of the present invention, a polypeptide, in particular an antigenic peptide, such as is defined above, or an anti-  
20 ligand, in particular an antibody to the said polypeptide;  
- all diagnostic, prophylactic or therapeutic compositions comprising one or more peptides, in particular antigenic peptides, such as are defined above, or one or more anti-ligands, in particular antibodies to the peptides,  
25 discussed above; such a composition is preferably, and by way of example, a vaccine composition.

The invention also relates to any diagnostic, prophylactic or therapeutic composition, in particular for inhibiting the expression of at least one virus associated 30 with MS or RA, and/or the enzymatic activities of the proteins of said virus, comprising a nucleotide fragment such as is defined above or a polynucleotide, in particular oligonucleotide, whose sequence is partially identical to that of the said fragment, except for that of  
35 the fragment having the nucleotide sequence SEQ ID NO:1. Likewise, it relates to any diagnostic, prophylactic or

therapeutic composition, in particular for inhibiting the expression of at least one pathogenic and/or infective agent associated with RA, comprising a nucleotide fragment such as is defined above by reference to the pol and gag genes, and especially with respect to the sequences SEQ ID NOS 40, 51, 56, 59, 60, 61, 62 and 89.

According to the invention, these same fragments or polynucleotides, in particular oligonucleotides, may participate in all suitable compositions for detecting, according to any suitable process or method, a pathological and/or infective agent associated with MS and with RA, respectively, in a biological sample. In such a process, an RNA and/or a DNA presumed to belong or originating from the said pathological and/or infective agent, and/or their complementary RNA and/or DNA, is/are brought into contact with such a composition.

The present invention also relates to any process for detecting the presence or exposure to such a pathological and/or infective agent, in a biological sample, by bringing this sample into contact with a peptide, in particular an antigenic peptide such as is defined above, or an anti-ligand, in particular an antibody to this peptide, such as is defined above.

In practice, and for example, a device for detection of the MSRV-1 virus comprises a reagent such as is defined above, supported by a solid support which is immunologically compatible with the reagent, and a means for bringing the biological sample, for example a sample of blood or of cerebrospinal fluid, likely to contain anti-MSRV-1 antibodies, into contact with this reagent under conditions permitting a possible immunological reaction, the foregoing items being accompanied by means for detecting the immune complex formed with this reagent.

Lastly, the invention also relates to the detection of anti-MSRV-1 antibodies in a biological sample, for example a sample of blood or of cerebrospinal fluid,

according to which this sample is brought into contact with a reagent such as is defined above, consisting of an antibody, under conditions permitting their possible immunological reaction, and the presence of the immune complex thereby formed with the reagent is then detected.

Before describing the invention in detail, different terms used in the description and the claims are now defined:

- strain or isolate is understood to mean any infective and/or pathogenic biological fraction containing, for example, viruses and/or bacteria and/or parasites, generating pathogenic and/or antigenic power, harboured by a culture or a living host; as an example, a viral strain according to the above definition can contain a coinfective agent, for example a pathogenic protist,

- the term "MSRV" used in the present description denotes any pathogenic and/or infective agent associated with MS, in particular a viral species, the attenuated strains of the said viral species or the defective-interfering particles or particles containing coencapsidated genomes, or alternatively genomes recombined with a portion of the MSRV-1 genome, derived from this species. Viruses, and especially viruses containing RNA, are known to have a variability resulting, in particular, from relatively high rates of spontaneous mutation (7), which will be borne in mind below for defining the notion of equivalence,

- human virus is understood to mean a virus capable of infecting, or of being harboured by human beings,

- in view of all the natural or induced variations and/or recombination which may be encountered when implementing the present invention, the subjects of the latter, defined above and in the claims, have been expressed including the equivalents or derivatives of the different biological materials defined below, in

particular of the homologous nucleotide or peptide sequences,

- the variant of a virus or of a pathogenic and/or infective agent according to the invention  
5 comprises at least one antigen recognized by at least one antibody directed against at least one corresponding antigen of the said virus and/or said pathogenic and/or infective agent, and/or a genome any part of which is detected by at least one hybridization probe and/or at  
10 least one nucleotide amplification primer specific for the said virus and/or pathogenic and/or infective agent, such as, for example, for the MSRV-1 virus, the primers and probes having a nucleotide sequence chosen from SEQ ID NO:20 to SEQ ID NO:24, SEQ ID NO:26, SEQ ID NO:16...  
15 to SEQ ID NO:19, SEQ ID NO:31 to SEQ ID NO:33, SEQ ID NO:45, SEQ ID NO:47, SEQ ID NO:48, SEQ ID NO:49, SEQ ID NO:50, SEQ ID NO:45 and their complementary sequences, under particular hybridization conditions well known to a person skilled in the art,

20 - according to the invention, a nucleotide fragment or an oligonucleotide or polynucleotide is an arrangement of monomers, or a biopolymer, characterized by the informational sequence of the natural nucleic acids, which is capable of hybridizing with any other nucleotide  
25 fragment under predetermined conditions, it being possible for the arrangement to contain monomers of different chemical structures and to be obtained from a molecule of natural nucleic acid and/or by genetic recombination and/or by chemical synthesis; a nucleotide fragment may be  
30 identical to a genomic fragment of the MSRV-1 virus discussed in the present invention, in particular a gene of this virus, for example pol or env in the case of the said virus,

- thus, a monomer can be a natural nucleotide of  
35 nucleic acid whose constituent elements are a sugar, a phosphate group and a nitrogenous base; in RNA the sugar

is ribose, in DNA the sugar is 2-deoxyribose; depending on whether the nucleic acid is DNA or RNA, the nitrogenous base is chosen from adenine, guanine, uracil, cytosine and thymine; or the nucleotide can be modified in at least one 5 of the three constituent elements; as an example, the modification can occur in the bases, generating modified bases such as inosine, 5-methyldeoxycytidine, deoxyuridine, 5-(dimethylamino)deoxyuridine, 2,6-diaminopurine, 5-bromodeoxyuridine and any other modified 10 base promoting hybridization; in the sugar, the modification can consist of the replacement of at least one deoxyribose by a polyamide (8), and in the phosphate group, the modification can consist of its replacement by esters chosen, in particular, from diphosphate, alkyl- and 15 arylphosphonate and phosphorothioate esters,

- "informational sequence" is understood to mean any ordered succession of monomers whose chemical nature and order in a reference direction constitute or otherwise an item of functional information of the same quality as 20 that of the natural nucleic acids,

- hybridization is understood to mean the process during which, under suitable working conditions, two nucleotide fragments having sufficiently complementary sequences pair to form a complex structure, in particular 25 double or triple, preferably in the form of a helix,

- a probe comprises a nucleotide fragment synthesized chemically or obtained by digestion or enzymatic cleavage of a longer nucleotide fragment, comprising at least six monomers, advantageously from 10 to 1000 monomers, preferably 10 to 30 monomers and more preferably 18 30 to 30, and possessing a specificity of hybridization under particular conditions; preferably, a probe possessing fewer than 10 monomers, but preferably fewer than 15 monomers is not used alone, but is used in the presence of 35 other probes of equally short size or otherwise; under certain special conditions, it may be useful to use probes

of size greater than 100 monomers; a probe may be used, in particular, for diagnostic purposes, such molecules being, for example, capture and/or detection probes,

5 - the capture probe may be immobilized on a solid support by any suitable means, that is to say directly or indirectly, for example by covalent bonding or passive adsorption,

10 - the detection probe may be labelled by means of a label chosen, in particular, from radioactive isotopes, enzymes chosen, in particular, from peroxidase and alkaline phosphatase and those capable of hydrolysing a chromogenic, fluorogenic or luminescent substrate, chromophoric chemical compounds, chromogenic, fluorogenic or luminescent compounds, nucleotide base analogues and 15 biotin,

- the probes used for diagnostic purposes of the invention may be employed in all known hybridization techniques, and in particular the techniques termed "DOT-BLOT" (9), "SOUTHERN BLOT" (10), "NORTHERN BLOT", which is 20 a technique identical to the "SOUTHERN BLOT" technique but which uses RNA as target, and the SANDWICH technique (11); advantageously, the SANDWICH technique is used in the present invention, comprising a specific capture probe and/or a specific detection probe, on the understanding 25 that the capture probe and the detection probe must possess an at least partially different nucleotide sequence,

- any probe according to the present invention can hybridize in vivo or in vitro with RNA and/or with DNA 30 in order to block the phenomena of replication, in particular translation and/or transcription, and/or to degrade the said DNA and/or RNA,

- a primer is a probe comprising at least six monomers, and advantageously from 10 to 30 monomers, and 35 preferably from 18 to 25 monomers, possessing a specificity of hybridization under particular conditions

for the initiation of an enzymatic polymerization, for example in an amplification technique such as PCR (polymerase chain reaction), in an elongation process such as sequencing, in a method of reverse transcription or the 5 like,

- two nucleotide or peptide sequences are termed equivalent or derived with respect to one another, or with respect to a reference sequence, if functionally the corresponding biopolymers can perform substantially the 10 same role, without being identical, as regards the application or use in question, or in the technique in which they participate; two sequences are, in particular, equivalent if they are obtained as a result of natural variability, in particular spontaneous mutation of the 15 species from which they have been identified, or induced variability, as are two homologous sequences, homology being defined below,

- "variability" is understood to mean any spontaneous or induced modification of a sequence, in particular by substitution and/or insertion and/or deletion 20 of nucleotides and/or of nucleotide fragments, and/or extension and/or shortening of the sequence at one or both ends; an unnatural variability can result from the genetic engineering techniques used, for example the choice of 25 synthesis primers, degenerate or otherwise, selected for amplifying a nucleic acid; this variability can manifest itself in modifications of any starting sequence, considered as reference, and capable of being expressed by a degree of homology relative to the said reference 30 sequence,

- homology characterizes the degree of identity of two nucleotide or peptide fragments compared; it is measured by the percentage identity which is determined, in particular, by direct comparison of nucleotide or 35 peptide sequences, relative to reference nucleotide or peptide sequences,

- this percentage identity has been specifically determined for the nucleotide fragments, clones in particular, dealt with in the present invention, which are homologous to the fragments identified, for the MSRV-1 virus, by SEQ ID NO:1 to NO:9, SEQ ID NO:46, SEQ ID NO:51 to SEQ ID NO:53, SEQ ID NO:40, SEQ ID NO:56, SEQ ID NO:57 and SEQ ID NO:93, as well as for the probes and primers homologous to the probes and primers identified by SEQ ID NO:20 to SEQ ID NO:24, SEQ ID NO:26, SEQ ID NO:16 to SEQ ID NO:19, SEQ ID NO:31 to SEQ ID NO:33, SEQ ID NO:45, SEQ ID NO:47, SEQ ID NO:48, SEQ ID NO:49, SEQ ID NO:50, SEQ ID NO:55, SEQ ID NO:40, SEQ ID NO:56, SEQ ID NO:57 and SEQ ID NO:99 to SEQ ID NO:111; as an example, the smallest percentage identity observed between the different general consensus sequences of nucleic acids obtained from fragments of MSRV-1 viral RNA, originating from the LM7PC and PLI-2 lines according to a protocol detailed later, is 67% in the region described in Figure 1,

- any nucleotide fragment is termed equivalent or derived from a reference fragment if it possesses a nucleotide sequence equivalent to the sequence of the reference fragment; according to the above definition, the following in particular are equivalent to a reference nucleotide fragment:

- 25 a) any fragment capable of hybridizing at least partially with the complement of the reference fragment,
- b) any fragment whose alignment with the reference fragment results in the demonstration of a larger number of identical contiguous bases than with any other fragment originating from another taxonomic group,
- c) any fragment resulting, or capable of resulting, from the natural variability of the species from which it is obtained,
- d) any fragment capable of resulting from the 35 genetic engineering techniques applied to the reference fragment,

e) any fragment containing at least eight contiguous nucleotides encoding a peptide which is homologous or identical to the peptide encoded by the reference fragment,

5 f) any fragment which is different from the reference fragment by insertion, deletion or substitution of at least one monomer, or extension or shortening at one or both of its ends; for example, any fragment corresponding to the reference fragment flanked at one or  
10 both of its ends by a nucleotide sequence not coding for a polypeptide,

- polypeptide is understood to mean, in particular, any peptide of at least two amino acids, in particular an oligopeptide, or protein, and for example an  
15 enzyme, extracted, separated or substantially isolated or synthesized through human intervention, in particular those obtained by chemical synthesis or by expression in a recombinant organism,

- polypeptide partially encoded by a nucleotide  
20 fragment is understood to mean a polypeptide possessing at least three amino acids encoded by at least nine contiguous monomers lying within the said nucleotide fragment,

- an amino acid is termed analogous to another  
25 amino acid when their respective physicochemical properties, such as polarity, hydrophobicity and/or basicity and/or acidity and/or neutrality are substantially the same; thus, a leucine is analogous to an isoleucine.

- any polypeptide is termed equivalent or  
30 derived from a reference polypeptide if the polypeptides compared have substantially the same properties, and in particular the same antigenic, immunological, enzymological and/or molecular recognition properties; the following in particular are equivalent to a reference  
35 polypeptide:

a) any polypeptide possessing a sequence in which at least one amino acid has been replaced by an analogous amino acid,

5 b) any polypeptide having an equivalent peptide sequence, obtained by natural or induced variation of the said reference polypeptide and/or of the nucleotide fragment coding for the said polypeptide,

c) a mimotope of the said reference polypeptide,

10 d) any polypeptide in whose sequence one or more amino acids of the L series are replaced by an amino acid of the D series, and vice versa,

15 e) any polypeptide into whose sequence a modification of the side chains of the amino acids has been introduced, such as, for example, an acetylation of the amine functions, a carboxylation of the thiol functions, an esterification of the carboxyl functions,

20 f) any polypeptide in whose sequence one or more peptide bonds have been modified, such as, for example, carba, retro, inverso, retro-inverso, reduced and methyleneoxy bonds,

(g) any polypeptide at least one antigen of which is recognized by an antibody directed against a reference polypeptide,

25 - the percentage identity characterizing the homology of two peptide fragments compared is, according to the present invention, at least 50% and preferably at least 70%.

In view of the fact that a virus possessing reverse transcriptase enzymatic activity may be genetically characterized equally well in RNA and in DNA form, both the viral DNA and RNA will be referred to for characterizing the sequences relating to a virus possessing such reverse transcriptase activity, termed MSRV-1 according to the present description.

35 The expressions of order used in the present description and the claims, such as "first nucleotide

sequence", are not adopted so as to express a particular order, but so as to define the invention more clearly.

Detection of a substance or agent is understood below to mean both an identification and a quantification, 5 or a separation or isolation, of the said substance or said agent.

A better understanding of the invention will be gained on reading the detailed description which follows, prepared with reference to the attached figures, in which:

10 - Figure 1 shows general consensus sequences of nucleic acids of the MSRV-1B clones amplified by the PCR technique in the "pol" region defined by Shih (12), from viral DNA originating from the LM7PC and PLI-2 lines, and identified under the references SEQ ID NO:3, SEQ ID NO:4, 15 SEQ ID NO:5 and SEQ ID NO:6, and the common consensus with amplification primers bearing the reference SEQ ID NO:7;

20 - Figure 2 gives the definition of a functional reading frame for each MSRV-1B/"PCR pol" type family, the said families A to D being defined, respectively, by the nucleotide sequences SEQ ID NO:3, SEQ ID NO:4, SEQ ID NO:5 and SEQ ID NO:6 described in Figure 1;

- Figure 3 gives an example of consensus of the MSRV-2B sequences, identified by SEQ ID NO:11;

25 - Figure 4 is a representation of the reverse transcriptase (RT) activity in dpm (disintegrations per minute) in the sucrose fractions taken from a purification gradient of the virions produced by the B lymphocytes in culture from a patient suffering from MS;

30 - Figure 5 gives, under the same experimental conditions as in Figure 4, the assay of the reverse transcriptase activity in the culture of a B lymphocyte line obtained from a control free from MS;

- Figure 6 shows the nucleotide sequence of the clone PSJ17 (SEQ ID NO:9);

35 - Figure 7 shows the nucleotide sequence SEQ ID NO:8 of the clone designated M003-P004;

- Figure 8 shows the nucleotide sequence SEQ ID NO:2 of the clone F11-1; the portion located between the two arrows in the region of the primer corresponds to a variability imposed by the choice of primer which was used 5 for the cloning of F11-1; in this same figure, the translation into amino acids is shown;

- Figure 9 shows the nucleotide sequence SEQ ID NO:1, and a possible functional reading frame of SEQ ID NO:1 in terms of amino acids; on this sequence, the 10 consensus sequences of the pol gene are underlined;

- Figures 10 and 11 give the results of a PCR, in the form of a photograph under ultraviolet light of an ethidium bromide-impregnated agarose gel, of the amplification products obtained from the primers identified by 15 SEQ ID NO:16, SEQ ID NO:17, SEQ ID NO:18 and SEQ ID NO:19;

- Figure 12 gives a representation in matrix form of the homology between SEQ ID NO:1 of MSRV-1 and that of an endogenous retrovirus designated HSERV9; this homology of at least 65% is demonstrated by a continuous 20 line, the absence of a line meaning a homology of less than 65%;

- Figure 13 shows the nucleotide sequence SEQ ID NO:46 of the clone FBd3;

- Figure 14 shows the sequence homology between 25 the clone FBd3 and the HSERV-9 retrovirus;

- Figure 15 shows the nucleotide sequence SEQ ID NO:51 of the clone t pol;

- Figures 16 and 17 show, respectively, the nucleotide sequences SEQ ID NO:52 and SEQ ID NO:53 of the 30 clones JLBC1 and JLBC2, respectively;

- Figure 18 shows the sequence homology between the clone JLBC1 and the clone FBd3;

- and Figure 19 the sequence homology between the clone JLBC2 and the clone FBd3;

35 - Figure 20 shows the sequence homology between the clones JLBC1 and JLBC2;

- Figures 21 and 22 show the sequence homology between the HSERV-9 retrovirus and the clones JLBC1 and JLBC2, respectively;

5 - Figure 23 shows the nucleotide sequence SEQ ID NO:56 of the clone GM3;

- Figure 24 shows the sequence homology between the HSERV-9 retrovirus and the clone GM3;

10 - Figure 25 shows the localization of the different clones studied, relative to the genome of the known retrovirus ERV9;

- Figure 26 shows the position of the clones F11-1, M003-P004, MSRV-1B and PSJ17 in the region hereinafter designated MSRV-1 pol\*;

15 - Figure 27, split into three successive Figures 27a-27c, shows a possible reading frame covering the whole of the pol gene;

20 - Figure 28 shows, according to SEQ ID NO:40, the nucleotide sequence coding for the peptide fragment POL2B, having the amino acid sequence identified by SEQ ID NO:39;

- Figure 29 shows the OD values (ELISA tests) at 492 nm obtained for 29 sera of MS patients and 32 sera of healthy controls tested with an anti-IgG antibody;

25 - Figure 30 shows the OD values (ELISA tests) at 492 nm obtained for 36 sera of MS patients and 42 sera of healthy controls tested with an anti-IgM antibody;

30 - Figures 31 to 33 show the results obtained (relative intensity of the spots) for 43 overlapping octapeptides covering the amino acid sequence 61-110, according to the Spotscan technique, respectively with a pool of MS sera, with a pool of control sera and with the pool of MS sera after deduction of a background corresponding to the maximum signal detected on at least one octapeptide with the control serum (intensity = 1), on the understanding that these sera were diluted to 1/50. The

bar at the far right-hand end represents a graphic scale standard unrelated to the serological test;

- Figure 34 shows the SEQ ID NO:41 and SEQ ID NO:42 of two polypeptides comprising immunodominant 5 regions, while SEQ ID NO:43 and 44 represent immunoreactive polypeptides specific to MS;

- Figure 35 shows the nucleotide sequence SEQ ID NO:59 of the clone LB19 and three potential reading frames of SEQ ID NO:59 in terms of amino acids;

10 - Figure 36 shows the nucleotide sequence SEQ ID NO:88 (GAG\*) and a potential reading frame of SEQ ID NO:88 in terms of amino acids;

15 - Figure 37 shows the sequence homology between the clone FBd13 and the HSERV-9 retrovirus; according to this representation, the continuous line means a percentage homology greater than or equal to 70% and the absence of a line means a smaller percentage homology;

20 - Figure 38 shows the nucleotide sequence SEQ ID NO:61 of the clone FP6 and three potential reading frames of SEQ ID NO:61 in terms of amino acids;

- Figure 39 shows the nucleotide sequence SEQ ID NO:89 of the clone G+E+A and three potential reading frames of SEQ ID NO:89 in terms of amino acids;

25 - Figure 40 shows a reading frame found in the region E and coding for an MSRV-1 retroviral protease identified by SEQ ID NO:90;

30 - Figure 41 shows the response of each serum of patients suffering from MS, indicated by the symbol (+), and of healthy patients, symbolised by (-), tested with an anti-IgG antibody, expressed as net optical density at 492 nm;

35 - Figure 42 shows the response of each serum of patients suffering from MS, indicated by the symbols (+) and (QS), and of healthy patients (-), tested with an anti-IgM antibody, expressed as net optical density at 492 nm;

- Figure 43 shows the RT-activity profile in sucrose density gradients of pellets from B-cell lines supernatants; Control B-cell line ■ was obtained from the relative of a patient with mitochondriopathy. MS B-Cell 5 line □ was obtained from a patient with definite MS;

- Figure 44 shows the nucleotide and amino acid alignment of the conserved pol regions of viruses detected in the study (cf Example 18) by the "Pan-retrovirus" PCR. "Deletions" are represented by dashes and standard single-letter abbreviations are used to designate amino acids and nucleotides (i = inosine). The most highly conserved VLPQG and YXDD regions are shown as separate blocks in bold type at the end of each sequence. Amino acids which are present in all or in all but one of the sequences are underlined. 10 PCR primers (modified from (12)) PAN-UO and PAN-UI are orientated 5' to 3' (sense) whereas primer PAN-DI is 3' to 5' (antisense). Degeneracies are shown above (PAN-UO & PAN-DI) or below (PAN-UI) the PCR primer sequences. 15 "I" denotes the nine base 5' extension cttggatcc, "-I" denotes the nine base 5' extension ctcaagctt. The capture and detector probes DpV1 and CpV1b used in the ELOSA assay are shown below a representative MSRV-cpol sequence. At 20 three positions below the translated MSRV-cpol sequence alternative amino acids (representing "non-silent" nucleic acid variations) are shown in italics - K and Y substitutions were only observed in PLI-1 derived clones whereas R and W were encoded by a significant proportion of the clones irrespective of derivation. Note that DpV1 is peroxidase labelled and that CpV1b may be biotinylated 25 at the 5' end if streptavidin coated plates are used. The name of each sequence is indicated at the left of the figure.

HTLV1: Human Leukaemia Virus type 1; HIV1: Human Immunodeficiency Virus type 1; MoMLV: Moloney-Murine 30 Leukaemia Virus; MPMV: Mason-Pfizer Monkey Virus. ERV9:

Endogenous Retrovirus 9. MSRV-cpol: Multiple Sclerosis associated RetroVirus conserved pol region.

- Figure 45 shows a phylogenetic tree which is based on the conserved amino acid region encoded by the 5 pol gene of MSRV and of representative endogenous and exogenous retroviruses and DNA viruses with reverse transcriptase. It was generated by the U.P.G.M.A. tree program of Geneworks® software.

HSRV: Human Spumaretrovirus. EIAV: Equine Infectious 10 Aenemia Virus. BLV: Bovine Leukaemia Virus. HIV1, HIV2: Human Immunodeficiency Viruses type 1 and 2. HTLV1 and HTLV2: Human Leukaemia Viruses type 1 and 2. F-MuLV: Friend-Murine Leukaemia Virus. MoMLV: Moloney-Murine Leukaemia Virus. BAEV: Baboon Endogenous Virus. GaLV/ 15 Gibbon Ape Leukaemia Virus. HUMER41: Human Endogenous Retroviral sequence, clone 41. IAP: Intracisternal A-type Particle. MPMV: Mason-Pfizer Monkey Virus. HERVK10: Human Endogenous Retrovirus K10. MMTV: Mouse Mammary tumour Virus. HSERV9 (ERV9 database sequence): Human sequence of 20 Endogenous Retrovirus 9. MSRV: Multiple Sclerosis associated RetroVirus. SIV: Simian Immunodeficiency Virus; RTLV-H: Reverse Transcriptase-Like Viral sequence H; SFV: Simian Foamy Virus; VISNA: Visna retrovirus; SIV1: Simian Immunodeficiency Virus type 1; SRV-2: Simian Retrovirus 25 type 2; SMRV-H: Squirrel Monkey Retrovirus H.

- Figure 46 shows the MSRV sequence in the Protease and Reverse-Transcriptase regions of the pol gene.

The aminoacid translation is aligned under the 30 corresponding nucleotide sequence. The region corresponding to the Protease ORF cloned in a recombinant vector and expressed in *E. coli*, is boxed. The regions corresponding to the A and B fragments amplified on plasma samples from MS patients are indicated by brackets. The 35 Reverse-Transcriptase (RT) and RNase H (RNH) region is boxed with dotted line. The highly conserved aminoacids

and/or active sites of enzyme activities of both PRT and RT (including RNH) are shown underlined.

- Figure 47A illustrates the specific detection of MSRV-pol RNA sequence by RT-PCR in the sucrose density fraction associated with RT-activity and in MS plasma ; Figure 47B shows the RT-activity profile on a sucrose density gradient obtained with extracellular virion pelleted from an MS choroid-plexus culture. The photograph below shows an agarose gel loaded with PCR products amplified from round 1 (ST1.1) RT-PCR products with the ST1.2 primer set. From left to right: water control 1 from RT-PCR step with ST1.1 set; water control 2 amplified from water control 1 with ST1.2 nested primers; Molecular weight markers; Fraction n°1 to 10 corresponding to the RT-activity profile shown above; Plasma samples C1 and C2 from healthy blood donors. Plasma samples MS1 and MS2 from two MS patients.

- Figure 48 shows an example of a variant and/or recombined sequence in the region of the pol gene defined by homology with the overlapping regions described in Figure 25, as GM3, MSRV-1 pol\*, t pol and FBd3.

- Figure 49 shows the nucleotide (Figure 49A) and amino acid (Figure 49B) alignments of the pol region between clones 1, 5 and 8 of the same patient (Experiment 46-7).

- Figure 50 shows the nucleotide (Figure 50A) and amino acid (Figure 50B) alignments of the pol region between clones 41, 43 and 42 of the same patient (Experiment 68-1).

- Figure 51 shows the nucleotide (Figure 51A) and amino acid (Figure 51B) alignments of the pol region between the consensus sequence (SEQ ID NO: 176) of clones 1, 5 and 8 of the same patient (Experiment 46-7) and SEQ ID NO:1, and between their corresponding peptide sequences.

- Figure 52 shows the nucleotide (Figure 52A) and amino acid (Figure 52B) alignments of the pol region between the consensus sequence (SEQ ID NO: 169) of clones 41, 43 and 42 of the same patient (Experiment 68-1) and SEQ ID NO:1, and between their corresponding peptide sequences.

- Figure 53 shows the nucleotide (Figure 53A) and amino acid (Figure 53B) alignments of the pol region between the consensus sequence (SEQ ID NO: 176) of clones 1, 5 and 8 of the same patient (Experiment 46-7) and the consensus sequence (SEQ ID NO: 169) of clones 41, 43 and 42 of the same patient (Experiment 68-1).

Table 5 (at the end of the description) shows the sequences obtained by RT-PCR with degenerate pol primers on sucrose density gradient fractions containing the peak of RT-activity or its negative control (cf Example 18) ; and

Table 6 (at the end of the description) shows the clinical data and results of MSRV-cpol detection by "Pan-retro" PCR with specific ELOSA assay, on CSF from MS and control patients (cf Example 18).

**EXAMPLE 1: OBTAINING CLONES DESIGNATED MSRV-1B AND MSRV-2B, DEFINING, RESPECTIVELY, A RETROVIRUS MSRV-1 AND A COINFECTIVE AGENT MSRV2, BY "NESTED" PCR AMPLIFICATION OF THE CONSERVED POL REGIONS OF RETROVIRUSES ON VIRION PREPARATIONS ORIGINATING FROM THE LM7PC AND PLI-2 LINES**

A PCR technique derived from the technique published by Shih (12) was used. This technique enables all trace of contaminant DNA to be removed by treating all the components of the reaction medium with DNase. It concomitantly makes it possible, by the use of different but overlapping primers in two successive series of PCR amplification cycles, to increase the chances of amplifying a cDNA synthesized from an amount of RNA which is

small at the outset and further reduced in the sample by the spurious action of the DNase on the RNA. In effect, the DNase is used under conditions of activity in excess which enable all trace of contaminant DNA to be removed  
5 before inactivation of this enzyme remaining in the sample by heating to 85°C for 10 minutes. This variant of the PCR technique described by Shih (12) was used on a cDNA synthesized from the nucleic acids of fractions of infective particles purified on a sucrose gradient  
10 according to the technique described by H. Perron (13) from the "POL-2" isolate (ECACC No. V92072202) produced by the PLI-2 line (ECACC No. 92072201) on the one hand, and from the MS7PG isolate (ECACC No. V93010816) produced by the LM7PC line (ECACC No. 93010817) on the other hand.  
15 These cultures were obtained according to the methods which formed the subject of the patent applications published under Nos WO 93/20188 and WO 93/20189.

After cloning the products amplified by this technique with the TA Cloning Kit® and analysis of the  
20 sequence using an Applied Biosystems model 373A Automatic Sequencer, the sequences were analysed using the Geneworks® software on the latest available version of the Genebank® data bank.

The sequences cloned and sequenced from these  
25 samples correspond, in particular, to two types of sequence: a first type of sequence, to be found in the majority of the clones (55% of the clones originating from the POL-2 isolates of the PLI-2 culture, and 67% of the clones originating from the MS7PG isolates of the LM7PC  
30 cultures), which corresponds to a family of "pol" sequences closely similar to, but different from, the endogenous human retrovirus designated ERV-9 or HSERV-9, and a second type of sequence which corresponds to sequences very strongly homologous to a sequence  
35 attributed to another infective and/or pathogenic agent designated MSRV-2.

The first type of sequence, representing the majority of the clones, consists of sequences whose variability enables four subfamilies of sequences to be defined. These subfamilies are sufficiently similar to one another for it to be possible to consider them to be quasi-species originating from the same retrovirus, as is well known for the HIV-1 retrovirus (14), or to be the outcome of interference with several endogenous proviruses coregulated in the producing cells. These more or less defective endogenous elements are sensitive to the same regulatory signals possibly generated by a replicative provirus, since they belong to the same family of endogenous retroviruses (15). This new family of endogenous retroviruses, or alternatively this new retroviral species from which the generation of quasi-species has been obtained in culture, and which contains a consensus of the sequences described below, is designated MSRV-1B.

Figure 1 presents the general consensus sequences of the sequences of the different MSRV-1B clones sequenced in this experiment, these sequences being identified, respectively, by SEQ ID NO:3, SEQ ID NO:4, SEQ ID NO:5 and SEQ ID NO:6. These sequences display a homology with respect to nucleic acids ranging from 70% to 88% with the HSERV9 sequence referenced X57147 and M37638 in the Genebank® data base. Four "consensus" nucleic acid sequences representative of different quasi-species of a possibly exogenous retrovirus MSRV-1B, or of different subfamilies of an endogenous retrovirus MSRV-1B, have been defined. These representative consensus sequences are presented in Figure 2, with the translation into amino acids. A functional reading frame exists for each subfamily of these MSRV-1B sequences, and it can be seen that the functional open reading frame corresponds in each instance to the amino acid sequence appearing on the second line under the nucleic acid sequence. The general

consensus of the MSRV-1B sequence, identified by SEQ ID NO:7 and obtained by this PCR technique in the "pol" region, is presented in Figure 1.

The second type of sequence representing the 5 majority of the clones sequenced is represented by the sequence MSRV-2B presented in Figure 3 and identified by SEQ ID NO:11. The differences observed in the sequences corresponding to the PCR primers are explained by the use of degenerate primers in mixture form used under different 10 technical conditions.

The MSRV-2B sequence (SEQ ID NO:11) is sufficiently divergent from the retroviral sequences already described in the data banks for it to be suggested that the sequence region in question belongs to a new infective 15 agent, designated MSRV-2. This infective agent would, in principle, on the basis of the analysis of the first sequences obtained, be related to a retrovirus but, in view of the technique used for obtaining this sequence, it could also be a DNA virus whose genome codes for an enzyme 20 which incidentally possesses reverse transcriptase activity, as is the case, for example, with the hepatitis B virus, HBV (12). Furthermore, the random nature of the degenerate primers used for this PCR amplification technique may very well have permitted, as a result of 25 unforeseen sequence homologies or of conserved sites in the gene for a related enzyme, the amplification of a nucleic acid originating from a prokaryotic or eukaryotic pathogenic and/or coinfective agent (protist).

30           **EXAMPLE 2: OBTAINING CLONES DESIGNATED MSRV-1B AND MSRV-2B, DEFINING A FAMILY MSRV-1 and MSRV-2, BY "NESTED" PCR AMPLIFICATION OF THE CONSERVED POL REGIONS OF RETROVIRUSES ON PREPARATIONS OF B LYMPHOCYTES FROM A NEW CASE OF MS**

35           The same PCR technique, modified according to the technique of Shih (12), was used to amplify and

sequence the RNA nucleic acid material present in a purified fraction of virions at the peak of "LM7-like" reverse transcriptase activity on a sucrose gradient according to the technique described by H. Perron (13), 5 and according to the protocols mentioned in Example 1, from a spontaneous lymphoblastoid line obtained by self-immortalization in culture of B lymphocytes from an MS patient who was seropositive for the Epstein-Barr virus (EBV), after setting up the blood lymphoid cells in 10 culture in a suitable culture medium containing a suitable concentration of cyclosporin A. A representation of the reverse transcriptase activity in the sucrose fractions taken from a purification gradient of the virions produced by this line is presented in Figure 4. Similarly, the 15 culture supernatants of a B line obtained under the same conditions from a control free from MS were treated under the same conditions, and the assay of reverse transcriptase activity in the sucrose gradient fractions proved negative throughout (background), and is presented in Figure 5. Fraction 3 of the gradient corresponding to 20 the MS B line and the same fraction without reverse transcriptase activity of the non-MS control gradient were analysed by the same RT-PCR technique as before, derived from Shih (12), followed by the same steps of cloning and 25 sequencing as described in Example 1.

It is particularly noteworthy that the MSRV-1 and MSRV-2 type sequences are to be found only in the material associated with a peak of "LM7-like" reverse transcriptase activity originating from the MS B lymphoblastoid line. These sequences were not to be found with the material from the control (non-MS) B lymphoblastoid line in 26 recombinant clones taken at random. Only Mo-MuLV type contaminant sequences, originating from the commercial reverse transcriptase used for the cDNA 35 synthesis step, and sequences without any particular retroviral analogy were to be found in this control, as a

result of the "consensus" amplification of homologous polymerase sequences which is produced by this PCR technique. Furthermore, the absence of a concentrated target which competes for the amplification reaction in 5 the control sample permits the amplification of dilute contaminants. The difference in results is manifestly highly significant (chi-squared, p<0.001).

EXAMPLE 3: OBTAINING A CLONE PSJ17, DEFINING A  
10 RETROVIRUS MSRV-1, BY REACTION OF ENDOGENOUS REVERSE  
TRANSCRIPTASE WITH A VIRION PREPARATION ORIGINATING FROM  
THE PLI-2 LINE

This approach is directed towards obtaining reverse-transcribed DNA sequences from the supposedly retroviral RNA in the isolate using the reverse transcriptase activity present in this same isolate. This reverse transcriptase activity can theoretically function only in the presence of a retroviral RNA linked to a primer tRNA or hybridized with short strands of DNA 20 already reverse-transcribed in the retroviral particles (16). Thus, the obtaining of specific retroviral sequences in a material contaminated with cellular nucleic acids was optimized according to these authors by means of the specific enzymatic amplification of the portions of viral 25 RNAs with a viral reverse transcriptase activity. To this end, the authors determined the particular physicochemical conditions under which this enzymatic activity of reverse transcription on RNAs contained in virions could be effective in vitro. These conditions correspond to the 30 technical description of the protocols presented below (endogenous RT reaction, purification, cloning and sequencing).

The molecular approach consisted in using a preparation of concentrated but unpurified virion obtained 35 from the culture supernatants of the PLI-2 line, prepared according to the following method: the culture

supernatants are collected twice weekly, precentrifuged at 10,000 rpm for 30 minutes to remove cell debris and then frozen at -80°C or used as they are for the following steps. The fresh or thawed supernatants are centrifuged on 5 a cushion of 30% glycerol-PBS at 100,000 g (or 30,000 rpm in a type 45 T LKB-HITACHI rotor) for 2 h at 4°C. After removal of the supernatant, the sedimented pellet is taken up in a small volume of PBS and constitutes the fraction of concentrated but unpurified virion. This concentrated 10 but unpurified viral sample was used to perform a so-called endogenous reverse transcription reaction, as described below.

A volume of 200 ml of virion purified according to the protocol described above, and containing a reverse 15 transcriptase activity of approximately 1-5 million dpm, is thawed at 37°C until a liquid phase appears, and then placed on ice. A 5-fold concentrated buffer was prepared with the following components: 500 mM Tris-HCl pH 8.2; 75 mM NaCl; 25 mM MgCl<sub>2</sub>; 75 mM DTT and 0.10% NP 40; 100 ml 20 of 5X buffer + 25 ml of a 100 mM solution of dATP + 25 ml of a 100 mM solution of dTTP + 25 ml of a 100 mM solution of dGTP + 25 ml of a 100 mM solution of dCTP + 100 ml of sterile distilled water + 200 ml of the virion suspension (RT activity of 5 million DPM) in PBS were mixed and 25 incubated at 42°C for 3 hours. After this incubation, the reaction mixture is added directly to a buffered phenol/chloroform/isoamyl alcohol mixture (Sigma ref. P 3803); the aqueous phase is collected and one volume of sterile distilled water is added to the organic phase to 30 re-extract the residual nucleic acid material. The collected aqueous phases are combined, and the nucleic acids contained are precipitated by adding 3M sodium acetate pH 5.2 to 1/10 volume + 2 volumes of ethanol + 1 ml of glycogen (Boehringer-Mannheim ref. 901 393) and 35 placing the sample at -20°C for 4 h or overnight at +4°C. The precipitate obtained after centrifugation is then

washed with 70% ethanol and resuspended in 60 ml of distilled water. The products of this reaction were then purified, cloned and sequenced according to the protocol which will now be described: blunt-ended DNAs with 5 unpaired adenines at the ends were generated: a "filling-in" reaction was first performed: 25 ml of the previously purified DNA solution were mixed with 2 ml of a 2.5 mM solution containing, in equimolar amounts, dATP + dGTP + dTTP + dCTP/1 ml of T4 DNA polymerase (Boehringer-Mannheim 10 ref. 1004 786) / 5 ml of 10X "incubation buffer for restriction enzyme" (Boehringer-Mannheim ref. 1417 975) / 1 ml of a 1% bovine serum albumin solution / 16 ml of sterile distilled water. This mixture was incubated for 20 minutes at 11°C. 50 ml of TE buffer and 1 ml of 15 glycogen (Boehringer-Mannheim ref. 901 393) were added thereto before extraction of the nucleic acids with phenol/chloroform/isoamyl alcohol (Sigma ref. P 3803) and precipitation with sodium acetate as described above. The DNA precipitated after centrifugation is resuspended in 20 10 ml of 10 mM Tris buffer pH 7.5. 5 ml of this suspension were then mixed with 20 ml of 5X Tag buffer, 20 ml of 5 mM dATP, 1 ml (5U) of Tag DNA polymerase (AmpliTaq™) and 54 ml of sterile distilled water. This mixture is incubated for 2 h at 75°C with a film of oil on the 25 surface of the solution. The DNA suspended in the aqueous solution drawn off under the film of oil after incubation is precipitated as described above and resuspended in 2 ml of sterile distilled water. The DNA obtained was inserted into a plasmid using the TA Cloning™ kit. The 2 ml of DNA 30 solution were mixed with 5 ml of sterile distilled water, 1 ml of a 10-fold concentrated ligation buffer "10X LIGATION BUFFER", 2 ml of "pCRT™ VECTOR" (25 ng/ml) and 1 ml of "TA DNA LIGASE". This mixture was incubated overnight at 12°C. The following steps were carried out 35 according to the instructions of the TA Cloning™ kit (British Biotechnology). At the end of the procedure, the

white colonies of recombinant bacteria (white) were picked out in order to be cultured and to permit extraction of the plasmids incorporated according to the so-called "miniprep" procedure (17). The plasmid preparation from 5 each recombinant colony was cut with a suitable restriction enzyme and analysed on agarose gel. Plasmids possessing an insert detected under UV light after staining the gel with ethidium bromide were selected for sequencing of the insert, after hybridization with a 10 primer complementary to the Sp6 promoter present on the cloning plasmid of the TA cloning™ kit. The reaction prior to sequencing was then performed according to the method recommended for the use of the sequencing kit "Prism ready reaction kit dye deoxyterminator cycle sequencing kit" 15 (Applied Biosystems, ref. 401384), and automatic sequencing was carried out with an Applied Biosystems "Automatic Sequencer, model 373 A" apparatus according to the manufacturer's instructions.

Discriminating analysis on the computerized data 20 banks of the sequences cloned from the DNA fragments present in the reaction mixture enabled a retroviral type sequence to be revealed. The corresponding clone PSJ17 was completely sequenced, and the sequence obtained, presented in Figure 6 and identified by SEQ ID NO:9, was analysed 25 using the "Geneworks®" software on the updated "Genebank™" data banks. An identical sequence already described could not be found by analysis of the data banks. Only a partial homology with some known retroviral elements was to be found. The most useful relative homology relates to an 30 endogenous retrovirus designated ERV-9, or HSERV-9, according to the references (18).

EXAMPLE 4: PCR AMPLIFICATION OF THE NUCLEIC ACID SEQUENCE CONTAINED BETWEEN THE 5' REGION DEFINED BY THE CLONE "POL MSRV-1B" AND THE 3' REGION DEFINED BY THE CLONE PSJ17

5 Five oligonucleotides, M001, M002-A, M003-BCD, P004 and P005, were defined in order to amplify the RNA originating from purified POL-2 virions. Control reactions were performed so as to check for the presence of contaminants (reaction with water). The amplification  
10 consists of an RT-PCR step according to the protocol described in Example 2, followed by a "nested" PCR according to the PCR protocol described in the document EP-A-0,569,272. In the first RT-PCR cycle, the primers M001 and P004 or P005 are used. In the second PCR cycle,  
15 the primers M002-A or M003-BCD and the primer P004 are used. The primers are positioned as follows:

M002-A  
M003-BCD  
M001 \_\_\_\_\_ P004 P005  
20 \_\_\_\_\_ \_\_\_\_\_  
\_\_\_\_\_ \_\_\_\_\_  
POL-2 \_\_\_\_\_ RNA  
<-----> \_\_\_\_\_ <----->  
pol MSRV-1B PSJ17  
25

Their composition is:

primer M001: GGTCITICCICAIGG (SEQ ID NO:20)  
primer M002-A: TTAGGGATAGCCCTCATCTCT (SEQ ID NO:21)  
primer M003-BCD: TCAGGGATAGCCCCCATCTAT (SEQ ID NO:22)  
30 primer P004: AACCCTTGCCACTACATCAATT (SEQ ID NO:23)  
primer P005: GCGTAAGGACTCCTAGAGCTATT (SEQ ID NO:24)

The "nested" amplification product obtained, and designated M003-P004, is presented in Figure 7, and corresponds to the sequence SEQ ID NO:8.

EXAMPLE 5: AMPLIFICATION AND CLONING OF A PORTION OF THE MSRV-1 RETROVIRAL GENOME USING A SEQUENCE ALREADY IDENTIFIED, IN A SAMPLE OF VIRUS PURIFIED AT THE PEAK OF REVERSE TRANSCRIPTASE ACTIVITY

5 A PCR technique derived from the technique published by Frohman (19) was used. The technique derived makes it possible, using a specific primer at the 3' end of the genome to be amplified, to elongate the sequence towards the 5' region of the genome to be analysed. This  
10 technical variant is described in the documentation of the firm "Clontech Laboratories Inc.", (Palo-Alto California, USA) supplied with its product "5'-AmpliFINDERTM RACE Kit", which was used on a fraction of virion purified as described above.

15 The specific 3' primers used in the kit protocol,  
for the synthesis of the cDNA and the PCR amplification  
are, respectively, complementary to the following MSRV-1  
sequences:

20 cDNA:TCATCCATGTACCGAAGG (SEQ ID NO:25)  
amplification :ATGGGGTTCCCAAGTTCCCT (SEQ ID NO:26)

The products originating from the PCR were obtained after purification on agarose gel according to conventional methods (17), and then resuspended in 10 ml of distilled water. Since one of the properties of Taq polymerase consists in adding an adenine at the 3' end of each of the two DNA strands, the DNA obtained was inserted directly into a plasmid using the TA Cloning™ kit (British Biotechnology). The 2 ml of DNA solution were mixed with 5 ml of sterile distilled water, 1 ml of a 10-fold concentrated ligation buffer "10X LIGATION BUFFER", 2 ml of "pCRT™ VECTOR" (25 ng/ml) and 1 ml of "TA DNA LIGASE". This mixture was incubated overnight at 12°C. The following steps were carried out according to the instructions of the TA Cloning™ kit (British Biotechnology). At the end of the procedure, the white

colonies of recombinant bacteria (white) were picked out in order to be cultured and to permit extraction of the plasmids incorporated according to the so-called "mini-prep" procedure (17). The plasmid preparation from each 5 recombinant colony was cut with a suitable restriction enzyme and analysed on agarose gel. Plasmids possessing an insert detected under UV light after staining the gel with ethidium bromide were selected for sequencing of the insert, after hybridization with a primer complementary to 10 the Sp6 promoter present on the cloning plasmid of the TA Cloning™ Kit. The reaction prior to sequencing was then performed according to the method recommended for the use of the sequencing kit "Prism ready reaction kit dye deoxyterminator cycle sequencing kit" (Applied Biosystems, 15 ref. 401384), and automatic sequencing was carried out with an Applied Biosystems "Automatic Sequencer model 373 A" apparatus according to the manufacturer's instructions.

This technique was applied first to two 20 fractions of virion purified as described below on sucrose from the "POL-2" isolate produced by the PLI-2 line on the one hand, and from the MS7PG isolate produced by the LM7PC line on the other hand. The culture supernatants are collected twice weekly, precentrifuged at 10,000 rpm for 25 30 minutes to remove cell debris and then frozen at -80°C or used as they are for the following steps. The fresh or thawed supernatants are centrifuged on a cushion of 30% glycerol-PBS at 100,000 g (or 30,000 rpm in a type 45 T LKB-HITACHI rotor) for 2 h at 4°C. After removal of the 30 supernatant, the sedimented pellet is taken up in a small volume of PBS and constitutes the fraction of concentrated but unpurified virions. The concentrated virus is then applied to a sucrose gradient in sterile PBS buffer (15 to 50% weight/weight) and ultracentrifuged at 35,000 rpm 35 (100,000 g) for 12 h at +4°C in a swing-out rotor. 10 fractions are collected, and 20 ml are withdrawn from

each fraction after homogenization to assay the reverse transcriptase activity therein according to the technique described by H. Perron (3). The fractions containing the peak of "LM7-like" RT activity are then diluted in sterile  
5 PBS buffer and ultracentrifuged for one hour at 35,000 rpm (100,000 g) to sediment the viral particles. The pellet of purified virion thereby obtained is then taken up in a small volume of a buffer which is appropriate for the extraction of RNA. The cDNA synthesis reaction mentioned  
10 above is carried out on this RNA extracted from purified extracellular virion. PCR amplification according to the technique mentioned above enabled the clone F1-11 to be obtained, whose sequence, identified by SEQ ID NO:2, is presented in Figure 8.

15 This clone makes it possible to define, with the different clones previously sequenced, a region of considerable length (1.2 kb) representative of the "pol" gene of the MSRV-1 retrovirus, as presented in Figure 9. This sequence, designated SEQ ID NO:1, is reconstituted  
20 from different clones overlapping one another at their ends, correcting the artefacts associated with the primers and with the amplification or cloning techniques which would artificially interrupt the reading frame of the whole. This sequence will be identified below under the  
25 designation "MSRV-1 pol\* region". Its degree of homology with the HSERV-9 sequence is shown in Figure 12.

In Figure 9, the potential reading frame with its translation into amino acids is presented below the nucleic acid sequence.

30

EXAMPLE 6: DETECTION OF SPECIFIC MSRV-1 and MSRV-2 SEQUENCES IN DIFFERENT SAMPLES OF PLASMA ORIGINATING FROM PATIENTS SUFFERING FROM MS OR FROM CONTROLS

35

A PCR technique was used to detect the MSRV-1 and MSRV-2 genomes in plasmas obtained after taking blood

samples from patients suffering from MS and from non-MS controls onto EDTA.

Extraction of the RNAs from plasma was performed according to the technique described by P. Chomzynski 5 (20), after adding one volume of buffer containing guanidinium thiocyanate to 1 ml of plasma stored frozen at -80°C after collection.

For MSRV-2, the PCR was performed under the same conditions and with the following primers:

- 10 - 5' primer, identified by SEQ ID NO:14  
5' GTAGTTCGATGTAGAAAGCG 3';  
- 3' primer, identified by SEQ ID NO:15  
5' GCATCCGGCAACTGCACG 3'.

However, similar results were also obtained with 15 the following PCR primers in two successive amplifications by "nested" PCR on samples of nucleic acids not treated with DNase.

The primers used for this first step of 40 cycles with a hybridization temperature of 48°C are the 20 following:

- 5' primer, identified by SEQ ID NO:27  
5' GCCGATATCACCCGCCATGG 3', corresponding to a 5' MSRV-2 PCR primer, for a first PCR on samples from patients,  
25 - 3' primer, identified by SEQ ID NO:28  
5' GCATCCGGCAACTGCACG 3', corresponding to a 3' MSRV-2 PCR primer, for a first PCR on samples from patients.

After this step, 10 ml of the amplification 30 product are taken and used to carry out a second, so-called "nested" PCR amplification with primers located within the region already amplified. This second step takes place over 35 cycles, with a primer hybridization ("annealing") temperature of 50°C. The reaction volume is 35 100 ml.

The primers used for this second step are the following:

- 5' primer, identified by SEQ ID NO:29  
5' CGCGATGCTGGTGGAGAGC 3', corresponding to a 5' MSRV-2 PCR primer, for a nested PCR on samples from patients,
- 3' primer, identified by SEQ ID NO:30  
5' TCTCCACTCCGAATATTCCG 3', corresponding to a 3' MSRV-2 PCR primer, for a nested PCR on samples from 10 patients.

For MSRV-1, the amplification was performed in two steps. Furthermore, the nucleic acid sample is treated beforehand with DNase, and a control PCR without RT (AMV reverse transcriptase) is performed on the two 15 amplification steps so as to verify that the RT-PCR amplification comes exclusively from the MSRV-1 RNA. In the event of a positive control without RT, the initial aliquot sample of RNA is again treated with DNase and amplified again.

20 The protocol for treatment with DNase lacking RNase activity is as follows: the extracted RNA is aliquoted in the presence of "RNase inhibitor" (Boehringer-Mannheim) in water treated with DEPC at a final concentration of 1 mg in 10 ml; to these 10 ml, 1 ml 25 of "RNase-free DNase" (Boehringer-Mannheim) and 1.2 ml of pH 5 buffer containing 0.1 M/l sodium acetate and 5 mM/l MgSO<sub>4</sub> is added; the mixture is incubated for 15 min at 20°C and brought to 95°C for 1.5 min in a "thermocycler".

30 The first MSRV-1 RT-PCR step is performed according to a variant of the RNA amplification method as described in Patent Application No. EP-A-0,569,272. In particular, the cDNA synthesis step is performed at 42°C for one hour; the PCR amplification takes place over 40 cycles, with a primer hybridization ("annealing") 35 temperature of 53°C. The reaction volume is 100 ml.

The primers used for this first step are the following:

- 5' primer, identified by SEQ ID NO:16  
5' AGGAGTAAGGAAACCCAACGGAC 3';
- 3' primer, identified by SEQ ID NO:17  
5' TAAGAGTTGCACAAGTGCG 3'.

5 After this step, 10 ml of the amplification product are taken and used to carry out a second, so-called "nested" PCR amplification with primers located 10 within the region already amplified. This second step takes place over 35 cycles, with a primer hybridization ("annealing") temperature of 53°C. The reaction volume is 100 ml.

15 The primers used for this second step are the following:

- 5' primer, identified by SEQ ID NO:18  
5' TCAGGGATAGCCCCCATCTAT 3';
- 3' primer, identified by SEQ ID NO:19  
5' AACCCCTTGCCACTACATCAATT 3'.

20 Figures 10 and 11 present the results of PCR in the form of photographs under ultraviolet light of ethidium bromide-impregnated agarose gels, in which an electrophoresis of the PCR amplification products applied separately to the different wells was performed.

25 The top photograph (Figure 10) shows the result of specific MSRV-2 amplification.

Well number 8 contains a mixture of DNA molecular weight markers, and wells 1 to 7 represent, in order, the products amplified from the total RNAs of 30 plasmas originating from 4 healthy controls free from MS (wells 1 to 4) and from 3 patients suffering from MS at different stages of the disease (wells 5 to 7).

In this series, MSRV-2 nucleic acid material is detected in the plasma of one case of MS out of the 35 tested, and in none of the 4 control plasmas. Other

results obtained on more extensive series confirm these results.

The bottom photograph (Figure 11) shows the result of specific amplification by MSRV-1 "nested"

5 RT-PCR:

well No. 1 contains the PCR product produced with water alone, without the addition of AMV reverse transcriptase; well No. 2 contains the PCR product produced with water alone, with the addition of AMV  
10 reverse transcriptase; well number 3 contains a mixture of DNA molecular weight markers; wells 4 to 13 contain, in order, the products amplified from the total RNAs extracted from sucrose gradient fractions (collected in a downward direction), on which gradient a pellet of virion  
15 originating from a supernatant of a culture infected with MSRV-1 and MSRV-2 was centrifuged to equilibrium according to the protocol described by H. Perron (13); to well 14 nothing was applied; to wells 15 to 17, the amplified products of RNA extracted from plasmas originating from  
20 3 different patients suffering from MS at different stages of the disease were applied.

The MSRV-1 retroviral genome is indeed to be found in the sucrose gradient fraction containing the peak of reverse transcriptase activity measured according to  
25 the technique described by H. Perron (3), with a very strong intensity (fraction 5 of the gradient, placed in well No. 8). A slight amplification has taken place in the first fraction (well No. 4), probably corresponding to RNA released by lysed particles which floated at the surface  
30 of the gradient; similarly, aggregated debris has sedimented in the last fraction (tube bottom), carrying with it a few copies of the MSRV-1 genome which have given rise to an amplification of low intensity.

Of the 3 MS plasmas tested in this series, MSRV-  
35 1 RNA turned up in one case, producing a very intense amplification (well No. 17).

In this series, the MSRV-1 retroviral RNA genome, probably corresponding to particles of extracellular virus present in the plasma in extremely small numbers, was detected by "nested" RT-PCR in one case 5 of MS out of the 3 tested. Other results obtained on more extensive series confirm these results.

Furthermore, the specificity of the sequences amplified by these PCR techniques may be verified and evaluated by the "ELOSA" technique as described by 10 F. Mallet (21) and in the document FR-A-2,663,040.

For MSRV-1, the products of the nested PCR described above may be tested in two ELOSA systems enabling a consensus A and a consensus B+C+D of MSRV-1 to be detected separately, corresponding to the subfamilies 15 described in Example 1 and Figures 1 and 2. In effect, the sequences closely resembling the consensus B+C+D are to be found essentially in the RNA samples originating from MSRV-1 virions purified from cultures or amplified in extracellular biological fluids of MS patients, whereas 20 the sequences closely resembling the consensus A are essentially to be found in normal human cellular DNA.

The ELOSA/MSRV-1 system for the capture and specific hybridization of the PCR products of the subfamily A uses a capture oligonucleotide cpV1A with an 25 amine bond at the 5' end and a biotinylated detection oligonucleotide dpV1A having as their sequence, respectively:

- cpV1A identified by SEQ ID NO:31
- 5' GATCTAGGCCACTTCTCAGGTCCAGS 3', corresponding 30 to the ELOSA capture oligonucleotide for the products of MSRV-1 nested PCR performed with the primers identified by SEQ ID NO:16 and SEQ ID NO:17, optionally followed by amplification with the primers identified by SEQ ID NO:18 and SEQ ID NO:19 on samples from patients;
- 35 - dpV1A identified by SEQ ID NO:32;

5' CATCTITTTGGICAGGCAITAGC 3', corresponding to the ELOSA capture oligonucleotide for the subfamily A of the products of MSRV-1 "nested" PCR performed with the primers identified by SEQ ID NO:16 and SEQ ID NO:17, 5 optionally followed by amplification with the primers identified by SEQ ID NO:18 and SEQ ID NO:19 on samples from patients.

The ELOSA/MSRV-1 system for the capture and specific hybridization of the PCR products of the 10 subfamily B+C+D uses the same biotinylated detection oligonucleotide dpV1A and a capture oligonucleotide cpV1B with an amine bond at the 5' end having as its sequence:

- dpV1B identified by SEQ ID NO:33

5' CTTGAGCCAGTTCTCATACCTGGA 3', corresponding to 15 the ELOSA capture oligonucleotide for the subfamily B + C + D of the products of MSRV-1 "nested" PCR performed with the primers identified by SEQ ID NO:16 and SEQ ID NO:17, optionally followed by amplification with the primers identified by SEQ ID NO:18 and SEQ ID NO:19 on samples 20 from patients.

This ELOSA detection system enabled it to be verified that none of the PCR products thus amplified from DNase-treated plasmas of MS patients contained a sequence of the subfamily A, and that all were positive with the 25 consensus of the subfamilies B, C and D.

For MSRV-2, a similar ELOSA technique was evaluated on isolates originating from infected cell cultures, using the following PCR amplification primers,

- 5' primer, identified by SEQ ID NO:34

30 5' AGTGYTRCCMCARGGCGCTGAA 3', corresponding to a 5' MSRV-2 PCR primer, for PCR on samples from cultures,

- 3' primer, identified by SEQ ID NO:35

5' GMGGGCCAGCAGSAKGTCATCCA 3', corresponding to a 3' MSRV-2 PCR primer, for PCR on samples from cultures,

and the capture oligonucleotides with an amine bond at the 5' end cpV2 and the biotinylated detection oligonucleotide dpV2 having as their respective sequences:

- cpV2 identified by SEQ ID NO:36

5 5' GGATGCCGCCCTATAAGCCTCTAC 3', corresponding to an ELOSA capture oligonucleotide for the products of MSRV-2 PCR performed with the primers SEQ ID NO:34 and SEQ ID NO:35, or optionally with the degenerate primers defined by Shih (12).

10 - dpV2 identified by SEQ ID NO:37

5' AAGCCTATCGCGTGCAGTTGCC 3', corresponding to an ELOSA detection oligonucleotide for the products of MSRV-2 PCR performed with the primers SEQ ID NO:34 and SEQ ID NO:35, or optionally with the degenerate primers 15 defined by Shih (12)

This PCR amplification system with a pair of primers different from those which were described previously for amplification on the samples from patients made it possible to confirm the infection with MSRV-2 of in 20 vitro cultures and of samples of nucleic acids used for the molecular biology studies.

All things considered, the first results of PCR detection of the genome of pathogenic and/or infective agents show that it is possible that free "virus" may 25 circulate in the blood stream of patients in an acute, virulent phase, outside the nervous system. This is compatible with the almost invariable presence of "gaps" in the blood-brain barrier of patients in an active phase of MS.

30

**EXAMPLE 7: OBTAINING SEQUENCES OF THE "env" GENE  
OF THE MSRV-1 RETROVIRAL GENOME**

As has already been described in Example 5, a 35 PCR technique derived from the technique published by Frohman (19) was used. The technique derived makes it possible, using a specific primer at the 3' end of the

genome to be amplified, to elongate the sequence towards the 5' region of the genome to be analysed. This technical variant is described in the documentation of "Clontech Laboratories Inc., (Palo-Alto California, USA) supplied 5 with its product "5'-AmpliFINDER™ RACE Kit", which was used on a fraction of virion purified as described above.

In order to carry out an amplification of the 3' region of the MSRV-1 retroviral genome encompassing the region of the "env" gene, a study was carried out to 10 determine a consensus sequence in the LTR regions of the same type as those of the defective endogenous retrovirus HSERV-9 (18, 24), with which the MSRV-1 retrovirus displays partial homologies.

The same specific 3' primer was used in the kit 15 protocol for the synthesis of the cDNA and the PCR amplification; its sequence is as follows:

GTGCTGATTGGTATTACAATCC (SEQ ID NO 45)

Synthesis of the complementary DNA (cDNA) and unidirectional PCR amplification with the above primer 20 were carried out in one step according to the method described in Patent EP-A-0,569,272.

The products originating from the PCR were extracted after purification of agarose gel according to conventional methods (17), and then resuspended in 10 ml 25 of distilled water. Since one of the properties of Taq polymerase consists in adding an adenine at the 3' end of each of the two DNA strands, the DNA obtained was inserted directly into a plasmid using the TA Cloning™ kit (British Biotechnology). The 2 ml of DNA solution were mixed with 5 30 ml of sterile distilled water, 1 ml of a 10-fold concentrated ligation buffer "10X LIGATION BUFFER", 2 ml of "pCRTM VECTOR" (25 ng/ml) and 1 ml of "TA DNA LIGASE". This mixture was incubated overnight at 12°C. The following steps were carried out according to the 35 instructions of the TA Cloning® kit (British Biotechnology). At the end of the procedure, the white colonies of

recombinant bacteria (white) were picked out in order to be cultured and to permit extraction of the plasmids incorporated according to the so-called "miniprep" procedure (17). The plasmid preparation from each 5 recombinant colony was cut with a suitable restriction enzyme and analysed on agarose gel. Plasmids possessing an insert detected under UV light after staining the gel with ethidium bromide were selected for sequencing of the insert, after hybridization with a primer complementary to 10 the Sp6 promoter present on the cloning plasmid of the TA Cloning™ Kit. The reaction prior to sequencing was then performed according to the method recommended for the use of the sequencing kit "Prism ready reaction kit dye deoxyterminator cycle sequencing kit" (Applied Biosystems, 15 ref. 401384), and automatic sequencing was carried out with an Applied Biosystems "automatic sequencer, model 373 A" apparatus according to the manufacturer's instructions.

This technical approach was applied to a sample 20 of virion concentrated as described below from a mixture of culture supernatants produced by B lymphoblastoid lines such as are described in Example 2, established from lymphocytes of patients suffering from MS and possessing reverse transcriptase activity which is detectable 25 according to the technique described by Perron et al. (3): the culture supernatants are collected twice weekly, precentrifuged at 10,000 rpm for 30 minutes to remove cell debris and then frozen at -80°C or used as they are for the following steps. The fresh or thawed supernatants are 30 centrifuged on a cushion of 30% glycerol-PBS at 100,000 g for 2 h at 4°C. After removal of the supernatant, the sedimented pellet constitutes the sample of concentrated but unpurified virions. The pellet thereby obtained is then taken up in a small volume of an appropriate buffer 35 for the extraction of RNA. The cDNA synthesis reaction

mentioned above is carried out on this RNA extracted from concentrated extracellular virion.

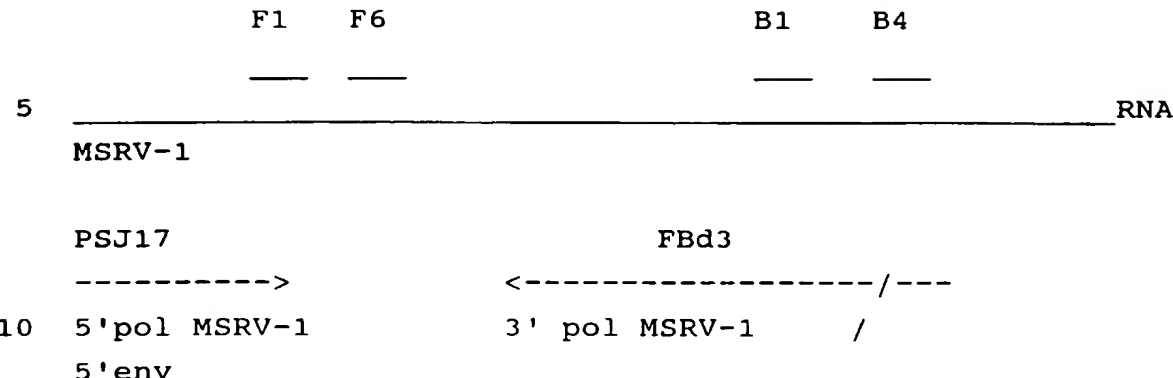
RT-PCR amplification according to the technique mentioned above enabled the clone FBd3 to be obtained, 5 whose sequence, identified by SEQ ID NO:46, is presented in Figure 13.

In Figure 14, the sequence homology between the clone FBd3 and the HSERV-9 retrovirus is shown on the matrix chart by a continuous line for any partial homology 10 greater than or equal to 65%. It can be seen that there are homologies in the flanking regions of the clone (with the pol gene at the 5' end and with the env gene and then the LTR at the 3' end), but that the internal region is totally divergent and does not display any homology, even 15 weak, with the "env" gene of HSERV9. Furthermore, it is apparent that the clone FBd3 contains a longer "env" region than the one which is described for the defective endogenous HSERV-9; it may thus be seen that the internal divergent region constitutes an "insert" between the 20 regions of partial homology with the HSERV-9 defective genes.

**EXAMPLE 8: AMPLIFICATION, CLONING AND SEQUENCING  
OF THE REGION OF THE MSRV-1 RETROVIRAL GENOME LOCATED  
25 BETWEEN THE CLONES PSJ17 AND FBd3**

Four oligonucleotides, F1, B4, F6 and B1, were defined for amplifying RNA originating from concentrated virions of the strains POL2 and MS7PG. Control reactions were performed so as to check for the presence of 30 contaminants (reaction with water). The amplification consists of a first step of RT-PCR according to the protocol described in Patent Application EP-A-0,569,272, followed by a second step of PCR performed on 10 ml of product of the first step with primers internal to the 35 amplified first region ("nested" PCR). In the first RT-PCR cycle, the primers F1 and B4 are used. In the second PCR

cycle, the primers F6 and the primer B1 are used. The primers are positioned as follows:



Their composition is:

primer F1: TGATGTGAAACGGCATACTCACTG (SEQ ID NO:47)  
15 primer B4: CCCAGAGGTTAGGAACCTCCCTTC (SEQ ID NO 48)  
primer F6: GCTAAAGGAGACTTGTGGTTGTCAG (SEQ ID NO 49)  
primer B1: CAACATGGGCATTTCGGATTAG (SEQ ID NO 50)

The product of "nested" amplification obtained  
and designated "t pol" is presented in Figure 15, and  
20 corresponds to the sequence SEQ ID NO:51.

EXAMPLE 9: OBTAINING NEW SEQUENCES, EXPRESSED AS RNA IN CELLS IN CULTURE PRODUCING MSRV-1, AND COMPRISING AN "env" REGION OF THE MSRV-1 RETROVIRAL GENOME

25 A library of cDNA was produced according to the  
procedure described by the manufacturer of the "cDNA  
synthesis module, cDNA rapid adaptor ligation module,  
cDNA rapid cloning module and lambda gt10 in vitro  
packaging module" kits (Amersham, ref RPN1256Y/Z, RPN1712,  
30 RPN1713, RPN1717, N334Z), from the messenger RNA extracted  
from cells of a B lymphoblastoid line such as is described  
in Example 2, established from the lymphocytes of a  
patient suffering from MS and possessing reverse  
transcriptase activity which is detectable according to  
35 the technique described by Perron et al. (3).

Oligonucleotides were defined for amplifying the cDNA cloned into the nucleic acid library between the 3' region of the clone PSJ17 (pol) and the 5'(LTR) region of the clone FBd3. Control reactions were performed so as to  
5 check for the presence of contaminants (reaction with water). PCR reactions performed on the nucleic acids cloned into the library with different pairs of primers enabled a series of clones linking pol sequences to the MSRV-1 type env or LTR sequences to be amplified.

10 Two clones are representative of the sequences obtained in the cellular cDNA library:

- the clone JLBC1, whose sequence SEQ ID NO:52 is presented in Figure 16;
- the clone JLBC2, whose sequence SEQ ID NO:53 is presented in Figure 17.

The sequences of the clones JLBC1 and JLBC2 are homologous to that of the clone FBd3, as is apparent in Figures 18 and 19. The homology between the clone JLBC1 and the clone JLBC2 is shown in Figure 20.

20 The homologies between the clones JLBC1 and JLBC2 on the one hand and the HSERV9 sequence on the other hand are presented, respectively, in Figures 21 and 22.

It will be noted that the region of homology between JLB1, JLB2 and FBd3 comprises, with a few sequence  
25 and size variations of the "insert", the additional sequence absent ("inserted") in the HSERV-9 env sequence, as described in Example 8.

It will also be noted that the cloned "pol" region is very homologous to HSERV-9, does not possess a  
30 reading frame (bearing in mind the sequence errors induced by the techniques used, including even the automatic sequencer) and diverges from the MSRV-1 sequences obtained from virions. In view of the fact that these sequences were cloned from the RNA of cells expressing MSRV-1  
35 particles, it is probable that they originate from endogenous retroviral elements related to the ERV9 family;

this is all the more likely for the fact that the pol and env genes are present on the same RNA which is clearly not the MSRV-1 genomic RNA. Some of these ERV9 elements possess functional LTRs which can be activated by 5 replicative viruses coding for homologous or heterologous transactivators. Under these conditions, the relationship between MSRV-1 and HSERV-9 makes probable the transactivation of the defective (or otherwise) endogenous ERV9 elements by homologous, or even identical, MSRV-1 10 transactivating proteins.

Such a phenomenon may induce a viral interference between the expression of MSRV-1 and the related endogenous elements. Such an interference generally leads to a so-called "defective-interfering" expression, some 15 features of which were to be found in the MSRV-1-infected cultures studied. Furthermore, such a phenomenon does not lack generation of the expression of polypeptides, or even of endogenous retroviral proteins which are not necessarily tolerated by the immune system. Such a scheme 20 of aberrant expression of endogenous elements related to MSRV-1 and induced by the latter is liable to multiply the aberrant antigens, and hence to contribute to the induction of autoimmune processes such as are observed in MS.

25 It is, however, essential to note that the clones JLBC1 and JLBC2 differ from the ERV9 or HSERV9 sequence already described, in that they possess a longer env region comprising an additional region totally divergent from ERV9. Their kinship with the endogenous 30 ERV9 family may hence be defined, but they clearly constitute novel elements never hitherto described. In effect, interrogation of the data banks of nucleic acid sequences available in version No. 15 (1995) of the "Entrez" software (NCBI, NIH, Bethesda, USA) did not 35 enable a known homologous sequence in the env region of these clones to be identified.

EXAMPLE 10: OBTAINING SEQUENCES LOCATED IN THE  
5' pol AND 3' gag REGION OF THE MSRV-1 RETROVIRAL GENOME

As has already been described in Example 5, a  
5 PCR technique derived from the technique published by  
Frohman (19) was used. The technique derived makes it  
possible, using a specific primer at the 3' end of the  
genome to be amplified, to elongate the sequence towards  
the 5' region of the genome to be analysed. This technical  
10 variant is described in the documentation of the firm  
Clontech Laboratories Inc., (Palo-Alto California, USA)  
supplied with its product "5'-AmpliFINDER™ RACE Kit",  
which was used on a fraction of virion purified as  
described above.

15 In order to carry out an amplification of the 5'  
region of the MSRV-1 retroviral genome starting from the  
pol sequence already sequenced (clone F11-1) and extending  
towards the gag gene, MSRV-1 specific primers were  
defined.

20 The specific 3' primers used in the kit protocol  
for the synthesis of the cDNA and the PCR amplification  
are, respectively, complementary to the following MSRV-1  
sequences:

cDNA: (SEQ ID NO:54)  
25 CCTGAGTTCTTGCACAAACCC  
amplification: (SEQ ID NO:55)  
GTCCCGTTGGGTTCCTTACTCCT

The products originating from the PCR were  
extracted after purification on agarose gel according to  
30 conventional methods (17), and then resuspended in 10 ml  
of distilled water. Since one of the properties of Taq  
polymerase consists in adding an adenine at the 3' end of  
each of the two DNA strands, the DNA obtained was inserted  
35 directly into a plasmid using the TA Cloning™ kit (British  
Biotechnology). The 2 ml of DNA solution were mixed with 5  
ml of sterile distilled water, 1 ml of a 10-fold

concentrated ligation buffer "10X LIGATION BUFFER", 2 ml of "pCRT™ VECTOR" (25 ng/ml) and 1 ml of "TA DNA LIGASE". This mixture was incubated overnight at 12°C. The following steps were carried out according to the 5 instructions of the TA Cloning® kit (British Biotechnology). At the end of the procedure, the white colonies of recombinant bacteria (white) were picked out in order to be cultured and to permit extraction of the plasmids incorporated according to the so-called 10 "miniprep" procedure (17). The plasmid preparation from each recombinant colony was cut with a suitable restriction enzyme and analysed on agarose gel. Plasmids possessing an insert detected under UV light after staining the gel with ethidium bromide were selected for 15 sequencing of the insert, after hybridization with a primer complementary to the Sp6 promoter present on the cloning plasmid of the TA Cloning™ Kit. The reaction prior to sequencing was then performed according to the method recommended for the use of the sequencing kit "Prism ready 20 reaction kit dye deoxyterminator cycle sequencing kit" (Applied Biosystems, ref. 401384), and automatic sequencing was carried out with an Applied Biosystems "automatic sequencer model 373 A" apparatus according to the manufacturer's instructions.

25 This technical approach was applied to a sample of virion concentrated as described below from a mixture of culture supernatants produced by B lymphoblastoid lines such as are described in Example 2, established from lymphocytes of patients suffering from MS and possessing 30 reverse transcriptase activity which is detectable according to the technique described by Perron et al. (3): the culture supernatants are collected twice weekly, precentrifuged at 10,000 rpm for 30 minutes to remove cell debris and then frozen at -80°C or used as they are for 35 the following steps. The fresh or thawed supernatants are centrifuged on a cushion of 30% glycerol-PBS at 100,000 g

for 2 h at 4°C. After removal of the supernatant, the sedimented pellet constitutes the sample of concentrated but unpurified virions. The pellet thereby obtained is then taken up in a small volume of an appropriate buffer 5 for the extraction of RNA. The cDNA synthesis reaction mentioned above is carried out on this RNA extracted from concentrated extracellular virion.

RT-PCR amplification according to the technique mentioned above enabled the clone GM3 to be obtained, 10 whose sequence, identified by SEQ ID NO 56, is presented in Figure 23.

In Figure 24, the sequence homology between the clone GMP3 and the HSERV-9 retrovirus is shown on the matrix chart by a continuous line, for any partial 15 homology greater than or equal to 65%.

In summary, Figure 25 shows the localization of the different clones studied above, relative to the known ERV9 genome. In Figure 25, since the MSRV-1 env region is longer than the reference ERV9 env gene, the additional 20 region is shown above the point of insertion according to a "V", on the understanding that the inserted material displays a sequence and size variability between the clones shown (JLBc1, JLBc2, FBd3). And Figure 26 shows the position of different clones studied in the MSRV-1 pol\* 25 region.

By means of the clone GM3 described above, a possible reading frame could be defined, covering the whole of the pol gene, referenced according to SEQ ID NO:57, shown in the successive Figures 27a to 27c.

30

**EXAMPLE 11: DETECTION OF ANTI-MSRV-1 SPECIFIC ANTIBODIES IN HUMAN SERUM**

Identification of the sequence of the pol gene of the MSRV-1 retrovirus and of an open reading frame of 35 this gene enabled the amino acid sequence SEQ ID NO:39 of

a region of the said gene, referenced SEQ ID NO:40, to be determined (see Figure 28).

Different synthetic peptides corresponding to fragments of the protein sequence of MSRV-1 reverse transcriptase encoded by the pol gene were tested for their antigenic specificity with respect to sera of patients suffering from MS and of healthy controls.

The peptides were synthesized chemically by solid-phase synthesis according to the Merrifield technique (Barany G, and Merrifield R.B, 1980, In the Peptides, 2, 1-284, Gross E and Meienhofer J, Eds., Academic Press, New York). The practical details are those described below.

a) Peptide synthesis:

The peptides were synthesized on a phenylacetamidomethyl (PAM)/polystyrene/divinylbenzene resin (Applied Biosystems, Inc. Foster City, CA), using an "Applied Biosystems 430A" automatic synthesizer. The amino acids are coupled in the form of hydroxybenzotriazole (HOBT) esters. The amino acids used are obtained from Novabiochem (Läuflelfingen, Switzerland) or Bachem (Bubendorf, Switzerland).

The chemical synthesis was performed using a double coupling protocol with N-methylpyrrolidone (NMP) as solvent. The peptides were cut from the resin, as well as the side-chain protective groups, simultaneously, using hydrofluoric acid (HF) in a suitable apparatus (type I cleavage apparatus, Peptide Institute, Osaka, Japan).

For 1 g of peptidyl resin, 10 ml of HF, 1 ml of anisole and 1 ml of dimethyl sulphide 5DMS are used. The mixture is stirred for 45 minutes at -2°C. The HF is then evaporated off under vacuum. After intensive washes with ether, the peptide is eluted from the resin with 10% acetic acid and then lyophilized.

The peptides are purified by preparative high performance liquid chromatography on a VYDAC C18 type

column (250 x 21 mm) (The Separation Group, Hesperia, CA, USA). Elution is carried out with an acetonitrile gradient at a flow rate of 22 ml/min. The fractions collected are monitored by an elution under isocratic conditions on a 5 VYDAC® C18 analytical column (250 x 4.6 mm) at a flow rate of 1 ml/min. Fractions having the same retention time are pooled and lyophilized. The preponderant fraction is then analysed by analytical high performance liquid chromatography with the system described above. The 10 peptide which is considered to be of acceptable purity manifests itself in a single peak representing not less than 95% of the chromatogram.

The purified peptides are then analysed with the object of monitoring their amino acid composition, using 15 an Applied Biosystems 420H automatic amino acid analyser. Measurement of the (average) chemical molecular mass of the peptides is obtained using LSIMS mass spectrometry in the positive ion mode on a VG. ZAB.ZSEQ double focusing instrument connected to a DEC-VAX 2000 acquisition system 20 (VG analytical Ltd, Manchester, England).

The reactivity of the different peptides was tested against sera of patients suffering from MS and against sera of healthy controls. This enabled a peptide designated POL2B to be selected, whose sequence is shown 25 in Figure 28 in the identifier SEQ ID NO:39, below, encoded by the pol gene of MSRV-1 (nucleotides 181 to 330).

b) Antigenic properties:

The antigenic properties of the POL2B peptide 30 were demonstrated according to the ELISA protocol described below.

The lyophilized POL2B peptide was dissolved in sterile distilled water at a concentration of 1 mg/ml. This stock solution was aliquoted and kept at +4°C for use 35 over a fortnight, or frozen at -20°C for use within 2 months. An aliquot is diluted in PBS (phosphate buffered

saline) solution so as to obtain a final peptide concentration of 1 microgram/ml. 100 microlitres of this dilution are placed in each well of microtitration plates ("high-binding" plastic, COSTAR ref: 3590). The plates are  
5 covered with a "plate-sealer" type adhesive and kept overnight at +4°C for the phase of adsorption of the peptide to the plastic. The adhesive is removed and the plates are washed three times with a volume of 300 micro-  
10 litres of a solution A (1X PBS, 0.05% Tween 20®), then inverted over an absorbent tissue. The plates thus drained are filled with 200 microlitres per well of a solution B (solution A + 10% of goat serum), then covered with an adhesive and incubated for 45 minutes to 1 hour at 37°C. The plates are then washed three times with the solution A  
15 as described above.

The test serum samples are diluted beforehand to 1/50 in the solution B, and 100 microlitres of each dilute test serum are placed in the wells of each microtitration plate. A negative control is placed in one well of each  
20 plate, in the form of 100 microlitres of buffer B. The plates covered with an adhesive are then incubated for 1 to 3 hours at 37°C. The plates are then washed three times with the solution A as described above. In parallel, a peroxidase-labelled goat antibody directed against human  
25 IgG (Sigma Immunochemicals ref. A6029) or IgM (Cappel ref. 55228) is diluted in the solution B (dilution 1/5000 for the anti-IgG and 1/1000 for the anti-IgM). 100 microlitres of the appropriate dilution of the labelled antibody are then placed in each well of the microtitration plates, and  
30 the plates covered with an adhesive are incubated for 1 to 2 hours at 37°C. A further washing of the plates is then performed as described above. In parallel, the peroxidase substrate is prepared according to the directions of the "Sigma fast OPD kit" (Sigma Immunochemicals, ref. P9187).  
35 100 microlitres of substrate solution are placed in each

well, and the plates are placed protected from light for 20 to 30 minutes at room temperature.

When the colour reaction has stabilized, the plates are placed immediately in an ELISA plate 5 spectrophotometric reader, and the optical density (OD) of each well is read at a wavelength of 492 nm. Alternatively, 30 microlitres of 1N HCl are placed in each well to stop the reaction, and the plates are read in the spectrophotometer within 24 hours.

10 The serological samples are introduced in duplicate or in triplicate, and the optical density (OD) corresponding to the serum tested is calculated by taking the mean of the OD values obtained for the same sample at the same dilution.

15 The net OD of each serum corresponds to the mean OD of the serum minus the mean OD of the negative control (solution B: PBS, 0.05% Tween 20®, 10% goat serum).

c) Detection of anti-MSRV-1 IgG antibodies by ELISA:

20 The technique described above was used with the POLB2 peptide to test for the presence of anti-MSRV-1 specific IgG antibodies in the serum of 29 patients for whom a definite or probable diagnosis of MS was established according to the criteria of Poser (23), and of 32 healthy controls (blood donors).

Figure 29 shows the results for each serum tested with an anti-IgG antibody. Each vertical bar represents the net optical density (OD at 492 nm) of a serum tested. The ordinate axis gives the net OD at the 30 top of the vertical bars. The first 29 vertical bars lying to the left of the vertical broken line represent the sera of 29 cases of MS tested, and the 32 vertical bars lying to the right of the vertical broken line represent the sera of 32 healthy controls (blood donors).

35 The mean of the net OD values for the MS sera tested is 0.62. The diagram enables 5 controls to be

revealed whose net OD rises above the grouped values of the control population. These values may represent the presence of specific IgGs in symptomless seropositive patients. Two methods were hence evaluated in order to 5 determine the statistical threshold of positivity of the test.

The mean of the net OD values for the controls, including the controls with high net OD values, is 0.36. Without the 5 controls whose net OD values are greater 10 than or equal to 0.5, the mean of the "negative" controls is 0.33. The standard deviation of the negative controls is 0.10. A theoretical threshold of positivity may be calculated according to the formula:  
threshold value (mean of the net OD values of the 15 seronegative controls) + (2 or 3 x standard deviation of the net OD values of the seronegative controls).

In the first case, there are considered to be symptomless seropositives, and the threshold value is equal to  $0.33 + (2 \times 0.10) = 0.53$ . The negative results 20 represent a non-specific "background" of the presence of antibodies directed specifically against an epitope of the peptide.

In the second case, if the set of controls consisting of blood donors in apparent good health is 25 taken as a reference basis, without excluding the sera which are, on the face of it, seropositive, the standard deviation of the "non-MS controls" is 0.116. The threshold value then becomes  $0.36 + (2 \times 0.116) = 0.59$ .

According to this analysis, the test is specific 30 for MS. In this respect, it is seen that the test is specific for MS, since, as shown in Table 1, no control has a net OD above this threshold. In fact, this result reflects the fact that the antibody titres in patients suffering from MS are, for the most part, higher than in 35 healthy controls who have been in contact with MSRV-1.

TABLE No. 1

	MS	CONTROLS
5	0.681	0.3515
	1.0425	0.56
	0.5675	0.3565
	0.63	0.449
	0.588	0.2825
10	0.645	0.55
	0.6635	0.52
	0.576	0.2535
	0.7765	0.55
	0.5745	0.51
15	0.513	0.426
	0.4325	0.451
	0.7255	0.227
	0.859	0.3905
	0.6435	0.265
20	0.5795	0.4295
	0.8655	0.291
	0.671	0.347
	0.596	0.4495
	0.662	0.3725
25	0.602	0.181
	0.525	0.2725
	0.53	0.426
	0.565	0.1915
	0.517	0.222
30	0.607	0.395
	0.3705	0.34
	0.397	0.307
	0.4395	0.219
		0.491
35		0.2265
		0.2605
	MEAN 0.62	0.33
	STD DEV 0.14	0.10
	THRESHOLD VALUE	0.53

In accordance with the first method of calculation, and as shown in Figure 29 and in the corresponding Table 1, 26 of the 29 MS sera give a positive result (net OD greater than or equal to 0.50), indicating the presence 5 of IgGs specifically directed against the POL2B peptide, hence against a portion of the reverse transcriptase enzyme of the MSRV-1 retrovirus encoded by its pol gene, and consequently against the MSRV-1 retrovirus. Thus, approximately 90% of the MS patients tested have reacted 10 against an epitope carried by the POL2B peptide and possess circulating IgGs directed against the latter.

Five out of 32 blood donors in apparent good health show a positive result. Thus, it is apparent that approximately 15% of the symptomless population may have 15 been in contact with an epitope carried by the POL2B peptide under conditions which have led to an active immunization which manifests itself in the persistence of specific serum IgGs. These conditions are compatible with an immunization against the MSRV-1 reverse 20 transcriptase during an infection with (and/or reactivation of) the MSRV-1 retrovirus. The absence of apparent neurological pathology recalling MS in these seropositive controls may indicate that they are healthy carriers and have eliminated an infectious virus after immunizing 25 themselves, or that they constitute an at-risk population of chronic carriers. In effect, epidemiological data showing that a pathogenic agent present in the environment of regions of high prevalence of MS may be the cause of this disease imply that a fraction of the population free 30 from MS has necessarily been in contact with such a pathogenic agent. It has been shown that the MSRV-1 retrovirus constitutes all or part of this "pathogenic agent" at the source of MS, and it is hence normal for controls taken from a healthy population to possess IgG 35 type antibodies against components of the MSRV-1 retrovirus. Thus, the difference in seroprevalence between

the MS and control populations is extremely significant: "chi-squared" test,  $p < 0.001$ . These results hence point to an aetiopathogenic role of MSRV-1 in MS.

d) Detection of anti-MSRV-1 IgM antibodies by  
5 ELISA:

The ELISA technique with the POL2B peptide was used to test for the presence of anti-MSRV-1 IgM specific antibodies in the serum of 36 patients for whom a definite or probable diagnosis of MS was established according to 10 the criteria of Poser (23), and of 42 healthy controls (blood donors).

Figure 30 shows the results for each serum tested with an anti-IgM antibody. Each vertical bar represents the net optical density (OD at 492 nm) of a serum tested. 15 The ordinate axis gives the net OD at the top of the vertical bars. The first 36 vertical bars lying to the left of the vertical line cutting the abscissa axis represent the sera of 36 cases of MS tested, and the vertical bars lying to the right of the vertical broken 20 line represent the sera of 42 healthy controls (blood donors). The horizontal line drawn in the middle of the diagram represents a theoretical threshold defining the boundary of the positive results (in which the top of the bar lies above) and the negative results (in which the top 25 of the bar lies below).

The mean of the net OD values for the MS cases tested is 0.19.

The mean of the net OD values for the controls is 0.09.

30 The standard deviation of the negative controls is 0.05.

In view of the small difference between the mean and the standard deviation of the controls, the threshold of theoretical positivity may be calculated according to 35 the formula:

threshold value = (mean of the net OD values of the seronegative controls) + (3 x standard deviation of the net OD values of the seronegative controls).

5           The threshold value is hence equal to 0.09 + (3 x 0.05) = 0.26; or, in practice, 0.25.

             The negative results represent a non-specific "background" of the presence of antibodies directed specifically against an epitope of the peptide.

10          According to this analysis, and as shown in Figure 30 and in the corresponding Table 2, the IgM test is specific for MS, since no control has a net OD above the threshold. 7 of the 36 MS sera produce a positive IgM result; now, a study of the clinical data reveals that 15 these positive sera were taken during a first attack of MS or an acute attack in untreated patients. It is known that IgMs directed against pathogenic agents are produced during primary infections or during reactivations following a latency phase of the said pathogenic agent.

20          The difference in seroprevalence between the MS and control populations is extremely significant: "chi-squared" test,  $p < 0.001$ .

             These results point to an aetiopathogenic role of MSRV-1 in MS.

25          The detection of IgM and IgG antibodies against the POL2B peptide enables the course of an MSRV-1 infection and/or of the viral reactivation of MSRV-1 to be evaluated.

TABLE No. 2

	MS	CONTROLS
5	0.064	0.243
	0.087	0.11
	0.044	0.098
	0.115	0.028
	0.089	0.094
	0.025	0.038
10	0.097	0.176
	0.108	0.146
	0.018	0.049
	0.234	0.161
	0.274	0.113
15	0.225	0.079
	0.314	0.093
	0.522	0.127
	0.306	0.02
	0.143	0.052
20	0.375	0.062
	0.142	0.074
	0.157	0.043
	0.168	0.046
	1.051	0.041
25	0.104	0.13
	0.187	0.153
	0.044	0.107
	0.053	0.178
	0.153	0.114
30	0.07	0.078
	0.033	0.118
	0.104	0.177
	0.187	0.026
	0.044	0.024
35	0.053	0.046
	0.153	0.116
	0.07	0.04
	0.033	0.028
	0.973	0.073
40		0.008
		0.074
		0.141
		0.219
		0.047
		0.017
	MEAN	0.19
45	STD. DEV.	0.23
	THRESHOLD VALUE	0.26

e) Search for immunodominant epitopes in the POL2B peptide:

In order to reduce the non-specific background and to optimize the detection of the responses of the 5 anti-MSRV-1 antibodies, the synthesis of octapeptides, advancing in successive one amino acid steps, covering the whole of the sequence determined by POL2B, was carried out according to the protocol described below.

The chemical synthesis of overlapping octapeptides 10 covering the amino acid sequence 61-110 shown in the identifier SEQ ID NO:39 was carried out on an activated cellulose membrane according to the technique of BERG et al. (1989. J. Ann. Chem. Soc., 111, 8024-8026) marketed by Cambridge Research Biochemicals under the trade name 15 Spotscan. This technique permits the simultaneous synthesis of a large number of peptides and their analysis.

The synthesis is carried out with esterified amino acids in which the α-amino group is protected with 20 an Fmoc group (Nova Biochem) and the side-chain groups with protective groups such as trityl, t-butyl ester or t-butyl ether. The esterified amino acids are solubilized in N-methylpyrrolidone (NMP) at a concentration of 300 nM, and 0.9 ml are applied to spots of deposit of bromophenol 25 blue. After incubation for 15 minutes, a further application of amino acids is carried out according to another 15-minute incubation. If the coupling between two amino acids has taken place correctly, a coloration modification (change from blue to yellow-green) is 30 observed. After three washes in DMF, an acetylation step is performed with acetic anhydride. Next, the terminal amino groups of the peptides in the process of synthesis are deprotected with 20% pyridine in DMF. The spots of deposit are restained with a 1% solution of bromophenol 35 blue in DMF, washed three times with methanol and dried. This set of operations constitutes one cycle of addition

of an amino acid, and this cycle is repeated until the synthesis is complete. When all the amino acids have been added, the NH<sub>2</sub>-terminal group of the last amino acid is deprotected with 20% piperidine in DMF and acetylated with 5 acetic anhydride. The groups protecting the side chain are removed with a dichloromethane/trifluoroacetic acid/triisobutylsilane (5 ml/5 ml/250 ml) mixture. The immunoreactivity of the peptides is then tested by ELISA.

After synthesis of the different octapeptides in 10 duplicate on two different membranes, the latter are rinsed with methanol and washed in TBS (0.1M Tris pH 7.2), then incubated overnight at room temperature in a saturation buffer. After several washes in TBS-T (0.1M Tris pH 7.2 - 0.05% Tween 20), one membrane is incubated 15 with a 1/50 dilution of a reference serum originating from a patient suffering from MS, and the other membrane with a 1/50 dilution of a pool of sera of healthy controls. The membranes are incubated for 4 hours at room temperature. After washes with TBS-T, a  $\beta$ -galactosidase-labelled anti- 20 human immunoglobulin conjugate (marketed by Cambridge Research Biochemicals) is added at a dilution of 1/200, and the mixture is incubated for two hours at room temperature. After washes of the membranes with 0.05% TBS-T and PBS, the immunoreactivity in the different spots is 25 visualized by adding 5-bromo-4-chloro-3-indolyl  $\beta$ -D-galactopyranoside in potassium. The intensity of coloration of the spots is estimated qualitatively with a relative value from 0 to 5 as shown in the attached Figures 31 to 33.

30 In this way, it is possible to determine two immunodominant regions at each end of the POL2B peptide, corresponding, respectively, to the amino acid sequences 65-75 (SEQ ID NO:41) and 92-109 (SEQ ID NO:42), according to Figure 34, and lying, respectively, between the 35 octapeptides Phe-Cys-Ile-Pro-Val-Arg-Pro-Asp (FCIPVRPD) and Arg-Pro-Asp-Ser-Gln-Phe-Leu-Phe (RPDSQFLF), and

Thr-Val-Leu-Pro-Gln-Gly-Phe-Arg (TVLPQGFR) and Leu-Phe-Gly-Gln-Ala-Leu-Ala-Gln (LFGQALAQ), and a region which is less reactive but apparently more specific, since it does not produce any background with the control serum,  
5 represented by the octapeptides Leu-Phe-Ala-Phe-Glu-Asp-Pro-Leu (LFAFEDPL) (SEQ ID NO:43) and Phe-Ala-Phe-Glu-Asp-Pro-Leu-Asn (FAFEDPLN) (SEQ ID NO:44).

These regions make it possible to define new peptides which are more specific and more immunoreactive  
10 according to the usual techniques.

It is thus possible, as a result of the discoveries made and the methods developed by the inventors, to carry out a diagnosis of MSRV-1 infection and/or reactivation and to evaluate a therapy in MS on the basis  
15 of its efficacy in "negativizing" the detection of these agents in the patients' biological fluids. Furthermore, early detection in individuals not yet displaying neurological signs of MS could make it possible to institute a treatment which would be all the more effective with  
20 respect to the subsequent clinical course for the fact that it would precede the lesion stage which corresponds to the onset of neurological disorders. Now, at the present time, a diagnosis of MS cannot be established before a symptomatology of neurological lesions has set  
25 in, and hence no treatment is instituted before the emergence of a clinical picture suggestive of lesions of the central nervous system which are already significant. The diagnosis of an MSRV-1 and/or MSRV-2 infection and/or reactivation in man is hence of decisive importance, and  
30 the present invention provides the means of doing this.

It is thus possible, apart from carrying out a diagnosis of MSRV-1 infection and/or reactivation, to evaluate a therapy in MS on the basis of its efficacy in "negativizing" the detection of these agents in the  
35 patients' biological fluids.

EXAMPLE 12: OBTAINING A CLONE LB19 CONTAINING A  
PORTION OF THE gag GENE OF THE MSRV-1 RETROVIRUS

A PCR technique derived from the technique published by Gonzalez-Quintial R et al. (19) and PLAZA et 5 al. (25) was used. From the total RNAs extracted from a fraction of virion purified as described above, the cDNA was synthesized using a specific primer (SEQ ID No.64) at the 3' end of the genome to be amplified, using EXPAND™ REVERSE TRANSCRIPTASE (BOEHRINGER MANNHEIM).

10

cDNA:

AAGGGGCATG      GACGAGGTGG      TGGCTTATT (SEQ      ID      NO:65)  
(antisense)

15

After purification, a poly(G) tail was added at the 5' end of the cDNA using the "Terminal transferases kit" marketed by the company Boehringer Mannheim, according to the manufacturer's protocol.

20

An anchoring PCR was carried out using the following 5' and 3' primers:

AGATCTGCAG    AATTCGATAT    CACCCCCCCC    CCCCCC (SEQ ID No. 91)  
(sense),      and      AAATGTCTGC      GGCACCAATC      TCCATGTT  
(SEQ ID No. 64) (antisense)

25

Next, a semi-nested anchoring PCR was carried out with the following 5' and 3' primers:

AGATCTGCAG    AATTCGATAT CA (SEQ ID No.92)    (sense),      and  
AAATGTCTGC    GGCACCAATC TCCATGTT (SEQ ID No.64) (antisense)

30

The products originating from the PCR were purified after purification on agarose gel according to conventional methods (17), and then resuspended in 10 microlitres of distilled water. Since one of the properties of Taq polymerase consists in adding an adenine at the 3' end of each of the two DNA strands, the DNA obtained was inserted directly into a plasmid using the TA 35 Cloning™ kit (British Biotechnology). The 2 µl of DNA solution were mixed with 5 µl of sterile distilled water,

1  $\mu$ l of 10-fold concentrated ligation buffer "10x LIGATION BUFFER", 2  $\mu$ l of "pCRTM VECTOR" (25 ng/ml) and 1  $\mu$ l of "T4 DNA LIGASE". This mixture was incubated overnight at 12°C. The following steps were carried out according to the 5 instructions of the TA CloningTM kit (British Biotechnology). At the end of the procedure, the white colonies of recombinant bacteria (white) were picked out in order to be cultured and to permit extraction of the plasmids incorporated according to the so-called 10 "miniprep" procedure (17). The plasmid preparation from each recombinant colony was cut with a suitable restriction enzyme and analysed on agarose gel. Plasmids possessing an insert detected under UV light after staining the gel with ethidium bromide were selected for 15 sequencing of the insert, after hybridization with a primer complementary to the Sp6 promoter present on the cloning plasmid of the TA Cloning KitTM. The reaction prior to sequencing was then performed according to the method recommended for the use of the sequencing kit "Prism ready 20 reaction kit dye deoxyterminator cycle sequencing kit" (Applied Biosystems, ref. 401384), and automatic sequencing was carried out with an Applied Biosystems "Automatic Sequencer, model 373 A" apparatus according to the manufacturer's instructions.

25 PCR amplification according to the technique mentioned above was used on a cDNA synthesized from the nucleic acids of fractions of infective particles purified on a sucrose gradient, according to the technique described by H. Perron (13), from culture supernatants of 30 B lymphocytes of a patient suffering from MS, immortalized with Epstein-Barr virus (EBV) strain B95 and expressing retroviral particles associated with reverse transcriptase activity as described by Perron et al. (3) and in French Patent Applications MS 10, 11 and 12. the clone LB19, 35 whose sequence, identified by SEQ ID NO:59, is presented in Figure 35.

The clone makes it possible to define, with the clone GM3 previously sequenced and the clone G+E+A (see Example 15), a region of 690 base pairs representative of a significant portion of the gag gene of the MSRV-1 retrovirus, as presented in Figure 36. This sequence designated SEQ ID NO:88 is reconstituted from different clones overlapping at their ends. This sequence is identified under the name MSRV-1 "gag\*" region. In Figure 36, a potential reading frame with the translation into 10 amino acids is presented below the nucleic acid sequence.

**EXAMPLE 13: OBTAINING A CLONE FBd13 CONTAINING A pol GENE REGION RELATED TO THE MSRV-1 RETROVIRUS AND AN APPARENTLY INCOMPLETE ENV REGION CONTAINING A POTENTIAL 15 READING FRAME (ORF) FOR A GLYCOPROTEIN**

Extraction of viral RNAs: The RNAs were extracted according to the method briefly described below.

A pool of culture supernatant of B lymphocytes of patients suffering from MS (650 ml) is centrifuged for 20 30 minutes at 10,000 g. The viral pellet obtained is resuspended in 300 microlitres of PBS/10 mM MgCl<sub>2</sub>. The material is treated with a DNase (100 mg/ml)/RNase (50 mg/ml) mixture for 30 minutes at 37°C and then with proteinase K (50 mg/ml) for 30 minutes at 46°C.

25 The nucleic acids are extracted with one volume of a phenol/0.1% SDS (V/V) mixture heated to 60°C, and then re-extracted with one volume of phenol/chloroform (1:1; V/V).

Precipitation of the material is performed with 30 2.5 V of ethanol in the presence of 0.1 V of sodium acetate pH5.2. The pellet obtained after centrifugation is resuspended in 50 microlitres of sterile DEPC water.

The sample is treated again with 50 mg/ml of "RNase free" DNase for 30 minutes at room temperature, 35 extracted with one volume of phenol/chloroform and

precipitated in the presence of sodium acetate and ethanol.

The RNA obtained is quantified by an OD reading at 260 nm. The presence of MSRV-1 and the absence of DNA 5 contaminant is monitored by a PCR and an MSRV-1-specific RTPCR associated with a specific ELOSA for the MSRV-1 genome.

Synthesis of cDNA:

5 mg of RNA are used to synthesize a cDNA primed 10 with a poly(DT) oligonucleotide according to the instructions of the "cDNA Synthesis Module" kit (ref RPN 1256, Amersham) with a few modifications: The reverse transcription is performed at 45°C instead of the recommended 42°C.

15 The synthesis product is purified by a double extraction and a double purification according to the manufacturer's instructions.

The presence of MSRV-1 is verified by an MSRV-1 PCR associated with a specific ELOSA for the MSRV-1 20 genome.

"Long Distance PCR": (LD-PCR)

500 ng of cDNA are used for the LD-PCR step (Expand Long Template System; Boehringer (ref.1681 842)).

Several pairs of oligonucleotides were used.

25 Among these, the pair defined by the following primers:

5' primer: GGAGAAAGAGC AGCATAAGTG G (SEQ ID NO:66)

3' primer: GTGCTGATTG GTGTATTTAC AATCC (SEQ ID NO:67).

The amplification conditions are as follows:

94°C 10 seconds

30 56°C 30 seconds

68°C 5 minutes;

10 cycles, then 20 cycles with an increment of 20 seconds in each cycle on the elongation time. At the end of this first amplification, 2 microlitres of the 35 amplification product are subjected to a second amplification under the same conditions as before.

The LD-PCR reactions are conducted in a Perkin model 9600 PCR apparatus in thin-walled microtubes (Boehringer).

5 The amplification products are monitored by electrophoresis of 1/5th of the amplification volume (10 microlitres) in 1% agarose gel. For the pair of primers described above, a band of approximately 1.7 Kb is obtained.

Cloning of the amplified fragment:

10 The PCR product was purified by passage through a preparative agarose gel and then through a Costar column (Spin; D. Dutcher) according to the supplier's instructions.

15 2 microlitres of the purified solution are joined up with 50 ng of vector PCRII according to the supplier's instructions (TA Cloning Kit; British Biotechnology)).

20 The recombinant vector obtained is isolated by transformation of competent DH5 $\alpha$ F' bacteria. The bacteria are selected using their resistance to ampicillin and the loss of metabolism for Xgal (= white colonies). The molecular structure of the recombinant vector is confirmed by plasmid minipreparation and hydrolysis with the enzyme EcoR1.

25 FBd13, a positive clone for all these criteria, was selected. A large-scale preparation of the recombinant plasmid was performed using the Midiprep Quiagen kit (ref 12243) according to the supplier's instructions.

30 Sequencing of the clone FBd13 is performed by means of the Perkin Prism Ready AmpliTaq FS dye terminator kit (ref. 402119) according to the manufacturer's instructions. The sequence reactions are introduced into a Perkin type 377 or 373A automatic sequencer. The sequencing strategy consists in gene walking carried out 35 on both strands of the clone Fbd13.

The sequence of the clone FBd13 is identified by SEQ ID NO:58.

In Figure 37, the sequence homology between the clone FBd13 and the HSERV-9 retrovirus is shown on the matrix chart by a continuous line for any partial homology greater than or equal to 70%. It can be seen that there are homologies in the flanking regions of the clone (with the pol gene at the 5' end and with the env gene and then the LTR at the 3' end), but that the internal region is totally divergent and does not display any homology, even weak, with the env gene of HSERV-9. Furthermore, it is apparent that the clone FBd13 contains a longer "env" region than the one which is described for the defective endogenous HSERV-9; it may thus be seen that the internal divergent region constitutes an "insert" between the regions of partial homology with the HSERV-9 defective genes.

This additional sequence determines a potential orf, designated ORF B13, which is represented by its amino acid sequence SEQ ID NO:87.

The molecular structure of the clone FBd13 was analyzed using the GeneWork software and Genebank and SwissProt data banks.

5 glycosylation sites were found.

The protein does not have significant homology with already known sequences.

It is probable that this clone originates from a recombination of an endogenous retroviral element (ERV), linked to the replication of MSRV-1.

Such a phenomenon does not lack generation of the expression of polypeptides, or even of endogenous retroviral proteins which are not necessarily tolerated by the immune system. Such a scheme of aberrant expression of endogenous elements related to MSRV-1 and/or induced by the latter is liable to multiply the aberrant antigens, and hence tends to contribute to the induction of

autoimmune processes such as are observed in MS. It clearly constitutes a novel element never hitherto described. In effect, interrogation of the data banks of nucleic acid sequences available in version No. 19 (1996) 5 of the "Entrez" software (NCBI, NIH, Bethesda, USA) did not enable a known homologous sequence comprising the whole of the env region of this clone to be identified.

EXAMPLE 14: OBTAINING A CLONE FP6 CONTAINING A  
10 PORTION OF THE pol GENE, WITH A REGION CODING FOR THE REVERSE TRANSCRIPTASE ENZYME HOMOLOGOUS TO THE CLONE POL\*  
MSRV-1, AND A 3'pol REGION DIVERGENT FROM THE EQUIVALENT SEQUENCES DESCRIBED IN THE CLONES POL\*, tpol, FBd3, JLBC1 and JLBC2

15 A 3'RACE was performed on total RNA extracted from plasma of a patient suffering from MS. A healthy control plasma treated under the same conditions was used as negative control. The synthesis of cDNA was carried out with the following modified oligo(dT) primer:

20 5' GACTCGCTGC AGATCGATT TTTTTTTTT TTTT 3' (SEQ ID NO:68)  
and Boehringer "Expand RT" reverse transcriptase according to the conditions recommended by the company. A PCR was performed with the enzyme Klentaq (Clontech) under the following conditions: 94°C 5 min then 93°C 1 min, 58°C 25 1 min, 68°C 3 min for 40 cycles and 68°C for 8 min, and with a final reaction volume of 50 µl.

Primers used for the PCR:

- 5' primer, identified by SEQ ID NO:69  
5' GCCATCAAGC CACCCAAGAA CTCTTAACCTT 3';  
30 - 3' primer, identified by SEQ ID NO:68 (=the same as for the cDNA)

A second, so-called "semi-nested" PCR was carried out with a 5' primer located within the region already amplified. This second PCR was performed under the 35 same experimental conditions as those used in the first

PCR, using 10 µl of the amplification product originating from the first PCR.

Primers used for the semi-nested PCR:

- 5' primer, identified by SEQ ID NO:70  
5 5' CCAATAGCCA GACCATTATA TACACTAATT 3';  
- 3' primer, identified by SEQ ID NO:68 (=the same as for the cDNA)

Primers SEQ ID NO:69 and SEQ ID NO:70 are specific for the pol\* region: position No. 403 to No. 422 10 and No. 641 to No. 670, respectively.

An amplification product was thus obtained from the extracellular RNA extracted from the plasma of a patient suffering from MS. The corresponding fragment was not observed for the plasma of the healthy control. This 15 amplification product was cloned in the following manner.

The amplified DNA was inserted into a plasmid using the TA Cloning™ kit. The 2 µl of DNA solution were mixed with 5 µl of sterile distilled water, 1 µl of a 10-fold concentrated ligation buffer "10x LIGATION 20 BUFFER", 2 µl of "PCR™ VECTOR" (25 ng/ml) and 1 µl of "TA DNA LIGASE". This mixture was incubated overnight at 12°C. The following steps were carried out according to the instructions of the TA Cloning™ kit (British Biotechnology). At the end of the procedure, the white 25 columns of recombinant bacteria (white) were picked out in order to be cultured and to permit extraction of the plasmids incorporated according to the so-called "miniprep" procedure (17). The plasmid preparation from each recombinant colony was cut with a suitable 30 restriction enzyme and analyzed on agarose gel. Plasmids possessing an insert detected under UV light after staining the gel with ethidium bromide was selected for sequencing of the insert, after hybridization with a 35 primer complementary to the Sp6 promoter present on the cloning plasmid of the TA cloning kit™. The reaction prior to sequencing was then performed according to the method

recommended for the use of the sequencing kit "Prism ready reaction kit dye deoxyterminator cycle sequencing kit" (Applied Biosystems, ref. 401384), and automatic sequencing was carried out with an Applied Biosystems 5 "Automatic Sequencer, model 373 A" apparatus according to the manufacturer's instructions.

The clone obtained, designated FP6, enables a region of 467 bp which is 89% homologous to the pol\* region of the MSRV-1 retrovirus and a region of 1167 bp 10 which is 64% homologous to the pol region of ERV-9 (No. 1634 to 2856) to be defined.

The clone FP6 is represented in Figure 38 by its nucleotide sequence identified by SEQ ID NO:61. The three potential reading frames of this clone are indicated by 15 their amino acid sequence under the nucleotide sequence.

EXAMPLE 15: OBTAINING A REGION DESIGNATED G+E+A  
CONTAINING AN ORF FOR A RETROVIRAL PROTEASE, BY PCR  
AMPLIFICATION OF THE NUCLEIC ACID SEQUENCE CONTAINED  
20 BETWEEN THE 5' REGION DEFINED BY THE CLONE "GM3" AND THE  
3' REGION DEFINED BY THE CLONE POL\*, FROM THE RNA  
EXTRACTED FROM A POOL OF PLASMAS OF PATIENTS SUFFERING  
FROM MS

Oligonucleotides specific for the MSRV-1 25 sequences already identified by the Applicant were defined in order to amplify the retroviral RNA originating from virions present in the plasma of patients suffering from MS. Control reactions were performed so as to monitor the presence of contaminants (reaction with water). The 30 amplification consists of a step of RT-PCR followed by a "nested" PCR. Pairs of primers were defined for amplifying three overlapping regions (designated G, E and A) on the regions defined by the sequences of the clones GM3 and pol\* described above.

- in the first RT-PCR cycle, the following primers are used:

primer 1: SEQ ID NO:71 (sense)

primer 2: SEQ ID NO:72 (antisense)

5 - in the second PCR cycle, the following primers  
are used:

primer 1: SEQ ID NO:73 (sense)

primer 4: SEQ ID NO:74 (antisense)

Nested RT-PCR for amplification of the region E:

10 - in the first RT-PCR cycle, the following primers are used:

primer 5: SEQ ID NO:75 (sense)

primer 6: SEQ ID NO:76 (antisense)

- in the second PCR cycle, the following primers  
15 are used:

primer 7: SEQ ID NO:77 (sense)

primer 8: SEQ ID NO:78 (antisense)

### Semi-nested RT-PCR for amplification of the region A:

- in the first RT-PCR cycle, the following  
20 primers are used:

primer 9: SEQ ID NO:79 (sense)

primer 10: SEQ ID NO:80 (antisense)

- in the second PCR cycle, the following primers are used:

25 primer 9: SEQ ID NO:81 (sense)  
primer 11: SEQ ID NO:82 (antisense)

The primers and the regions G, E and A which they define are positioned as follows:

cDNA

The sequence of the region defined by the different clones G, E and A was determined after cloning and sequencing of the "nested" amplification products.

The clones G, E and A were assembled together by 5 PCR with the primers 1 at the 5' end of the fragment G and 11 at the 3' end of the fragment A, the primers being described above. An approximately 1580-bp fragment G+E+A was amplified and inserted into a plasmid using the TA Cloning (trademark) kit. The sequence of the amplification 10 product corresponding to G+E+A was determined and analysis of the G+E and E+A overlaps was carried out. The sequence is shown in Figure 39, and corresponds to the sequence SEQ ID NO:89.

A reading frame coding for an MSRV-1 retroviral 15 protease was found in the region E. The amino acid sequence of the protease, identified by SEQ ID NO:90, is presented in Figure 40.

**EXAMPLE 16: OBTAINING A CLONE LTRGAG12, RELATED  
20 TO AN ENDOGENOUS RETROVIRAL ELEMENT (ERV) CLOSE TO MSRV-1,  
IN THE DNA OF AN MS LYMPHOBLASTOID LINE PRODUCING VIRIONS  
AND EXPRESSING THE MSRV-1 RETROVIRUS**

A nested PCR was performed on the DNA extracted from a lymphoblastoid line (B lymphocytes immortalized 25 with the EBV virus strain B95, as described above and as is well known to a person skilled in the art) expressing the MSRV-1 retrovirus and originating from peripheral blood lymphocytes of a patient suffering from MS.

In the first PCR step, the following primers are 30 used:

primer 4327: CTCGATTCT TGCTGGGCCT TA (SEQ ID NO:83)

primer 3512: GTTGATTCCTT TCCTCAAGCA (SEQ ID NO:84)

This step comprises 35 amplification cycles with the following conditions: 1 min at 94°C, 1 min at 54°C and 35 4 min at 72°C.

In the second PCR step, the following primers are used:

primer 4294: CTCTACCAAT CAGCATGTGG (SEQ ID NO:85)

primer 3591: TGTTCCCTCTT GGTCCCTAT (SEQ ID NO:86)

5 This step comprises 35 amplification cycles with the following conditions: 1 min at 94°C, 1 min at 54°C and 4 min at 72°C.

The products originating from the PCR were purified after purification on agarose gel according to 10 conventional methods (17), and then resuspended in 10 ml of distilled water. Since one of the properties of Taq polymerase consists in adding an adenine at the 3' end of each of the two DNA strands, the DNA obtained was inserted directly into a plasmid using the TA Cloning™ kit (British 15 Biotechnology). The 2 µl of DNA solution were mixed with 5 µl of sterile distilled water, 1 µl of a 10-fold concentrated ligation buffer "10x LIGATION BUFFER", 2 µl of "pCRTM VECTOR" (25 ng/ml) and 1 µl of "TA DNA LIGASE". This mixture was incubated overnight at 12°C. The 20 following steps were carried out according to the instructions of the TA Cloning™ kit (British Biotechnology). At the end of the procedure, the white colonies of recombinant bacteria (white) were picked out in order to be cultured and to permit extraction of the 25 plasmids incorporated according to the so-called "miniprep" procedure (17). The plasmid preparation from each recombinant colony was cut with a suitable restriction enzyme and analyzed on agarose gel. The plasmids possessing an insert detected under UV light 30 after staining the gel with ethidium bromide were selected for sequencing of the insert, after hybridization with a primer complementary to the Sp6 promoter present on the cloning plasmid of the TA Cloning Kit™. The reaction prior to sequencing was then performed according to the method 35 recommended for the use of the sequencing kit "Prism ready reaction kit dye deoxyterminator cycle sequencing kit"

(Applied Biosystems, ref. 401384), and automatic sequencing was carried out with an Applied Biosystems "Automatic Sequencer, model 373 A" apparatus according to the manufacturer's instructions.

5 Thus, a clone designated LTRGAG12 could be obtained, and is represented by its internal sequence identified by SEQ ID NO:60.

This clone is probably representative of endogenous elements close to ERV-9, present in human DNA, 10 in particular in the DNA of patients suffering from MS, and capable of interfering with the expression of the MSRV-1 retrovirus, hence capable of having a role in the pathogenesis associated with the MSRV-1 retrovirus and capable of serving as marker for a specific expression in 15 the pathology in question.

**EXAMPLE 17: DETECTION OF ANTI-MSRV-1 SPECIFIC ANTIBODIES IN HUMAN SERUM**

Identification of the sequence of the pol gene 20 of the MSRV-1 retrovirus and of an open reading frame of this gene enabled the amino acid sequence SEQ ID NO:63 of a region of the said gene, referenced SEQ ID NO:62, to be determined.

Different synthetic peptides corresponding to 25 fragments of the protein sequence of MSRV-1 reverse transcriptase encoded by the pol gene were tested for their antigenic specificity with respect to sera of patients suffering from MS and of healthy controls.

The peptides were synthesized chemically by 30 solid-phase synthesis according to the Merrifield technique (22). The practical details are those described below.

a) Peptide synthesis:

The peptides were synthesized on a phenylacet- 35 amidomethyl (PAM)/polystyrene/divinylbenzene resin (Applied Biosystems, Inc. Foster City, CA), using an

"Applied Biosystems 430A" automatic synthesizer. The amino acids are coupled in the form of hydroxybenzotriazole (HOBT) esters. The amino acids used are obtained from Novabiochem (Läuflelfingen, Switzerland) or Bachem 5 (Bubendorf, Switzerland).

The chemical synthesis was performed using a double coupling protocol with N-methylpyrrolidone (NMP) as solvent. The peptides were cut from the resin, as well as the side-chain protective groups, simultaneously, using 10 hydrofluoric acid (HF) in a suitable apparatus (type I cleavage apparatus, Peptide Institute, Osaka, Japan).

For 1 g of peptidyl resin, 10 ml of HF, 1 ml of anisole and 1 ml of dimethyl sulphide 5DMS are used. The mixture is stirred for 45 minutes at -2°C. The HF is then 15 evaporated off under vacuum. After intensive washes with ether, the peptide is eluted from the resin with 10% acetic acid and then lyophilized.

The peptides are purified by preparative high performance liquid chromatography on a VYDAC C18 type 20 column (250 x 21 mm) (The Separation Group, Hesperia, CA, USA). Elution is carried out with an acetonitrile gradient at a flow rate of 22 ml/min. The fractions collected are monitored by an elution under isocratic conditions on a VYDAC™ C18 analytical column (250 x 4.6 mm) at a flow rate 25 of 1 ml/min. Fractions having the same retention time are pooled and lyophilized. The preponderant fraction is then analysed by analytical high performance liquid chromatography with the system described above. The peptide which is considered to be of acceptable purity 30 manifests itself in a single peak representing not less than 95% of the chromatogram.

The purified peptides are then analysed with the object of monitoring their amino acid composition, using an Applied Biosystems 420H automatic amino acid analyser. 35 Measurement of the (average) chemical molecular mass of the peptides is obtained using LSIMS mass spectrometry in

the positive ion mode on a VG. ZAB.ZSEQ double focusing instrument connected to a DEC-VAX 2000 acquisition system (VG analytical Ltd, Manchester, England).

The reactivity of the different peptides was  
5 tested against sera of patients suffering from MS and against sera of healthy controls. This enabled a peptide designated S24Q to be selected, whose sequence is identified by SEQ ID NO:63, encoded by a nucleotide sequence of the pol gene of MSRV-1 (SEQ ID NO:62).

10

b) Antigenic properties:

The antigenic properties of the S24Q peptide were demonstrated according to the ELISA protocol described below.

15

The lyophilized S24Q peptide was dissolved in 10 % acetic acid at a concentration of 1 mg/ml. This stock solution was aliquoted and kept at +4°C for use over a fortnight, or frozen at -20°C for use within 2 months. An aliquot is diluted in PBS (phosphate buffered saline) 20 solution so as to obtain a final peptide concentration of 5 micrograms/ml. 100 microlitres of this dilution are placed in each well of Nunc Maxisorb (trade name) microtitration plates. The plates are covered with a "plate-sealer" type adhesive and kept for 2 hours at +37°C 25 for the phase of adsorption of the peptide to the plastic. The adhesive is removed and the plates are washed three times with a volume of 300 microlitres of a solution A (1X' PBS, 0.05% Tween 20®), then inverted over an absorbent tissue. The plates thus drained are filled with 30 250 microlitres per well of a solution B (solution A + 10% of goat serum), then covered with an adhesive and incubated for 1 hour at 37°C. The plates are then washed three times with the solution A as described above.

The test serum samples are diluted beforehand to 35 1/100 in the solution B, and 100 microlitres of each dilute test serum are placed in the wells of each micro-

titration plate. A negative control is placed in one well of each plate, in the form of 100 microlitres of buffer B. The plates covered with an adhesive are then incubated for 1 hour 30 min at 37°C. The plates are then washed three 5 times with the solution A as described above. For the IgG response, a peroxidase-labelled goat antibody directed against human IgG (marketed by Jackson Immuno Research Inc.) is diluted in the solution B (dilution 1/10,000). 100 microlitres of the appropriate dilution of the 10 labelled antibody are then placed in each well of the microtitration plates, and the plates covered with an adhesive are incubated for 1 hour at 37°C. A further washing of the plates is then performed as described above. In parallel, the peroxidase substrate is prepared 15 according to the directions of the bioMérieux kits. 100 microlitres of substrate solution are placed in each well, and the plates are placed protected from light for 20 to 30 minutes at room temperature.

When the colour reaction has stabilized, 20 50 microlitres of Color 2 (bioMérieux trade name) are placed in each well in order to stop the reaction. The plates are placed immediately in an ELISA plate spectrophotometric reader, and the optical density (OD) of each well is read at a wavelength of 492 nm.

25 The serological samples are introduced in duplicate or in triplicate, and the optical density (OD) corresponding to the serum tested is calculated by taking the mean of the OD values obtained for the same sample at the same dilution.

30 The net OD of each serum corresponds to the mean OD of the serum minus the mean OD of the negative control (solution B: PBS, 0.05% Tween 20x, 10% goat serum).

c) Detection of anti-MSRV-1 IgG antibodies (S24Q) by ELISA:

35 The technique described above was used with the S24Q peptide to test for the presence of anti-MSRV-1

specific IgG antibodies in the serum of 15 patients for whom a definite diagnosis of MS was established according to the criteria of Poser (23), and of 15 healthy controls (blood donors).

5         Figure 41 shows the results for each serum tested with an anti-IgG antibody. Each vertical bar represents the net optical density (OD at 492 nm) of a serum tested. The ordinate axis gives the net OD at the top of the vertical bars. The first 15 vertical bars lying 10 to the left of the vertical broken line represent the sera of 15 healthy controls (blood donors), and the 15 vertical bars lying to the right of the vertical broken line represent the sera of 15 cases of MS tested. The diagram enables 2 controls to be revealed whose OD rises above the 15 grouped values of the control population. These values may represent the presence of specific IgGs in symptomless seropositive patients. Two methods were hence evaluated in order to determine the statistical threshold of positivity of the test.

20         The mean of the net OD values for the controls, including the controls with high net OD values, is 0.129 and the standard deviation is 0.06. Without the 2 controls whose OD values are greater than 0.2, the mean of the "negative" controls is 0.107 and the standard deviation is 25 0.03. A theoretical threshold of positivity may be calculated according to the formula:

threshold value (mean of the net OD values of the negative controls) + ( 2 or 3 ' standard deviation 30 of the net OD values of the negative controls).

In the first case, there are considered to be 35 symptomless seropositives, and the threshold value is equal to  $0.11 + (3 \times 0.03) = 0.20$ . The negative results represent a non-specific "background" of the presence of

antibodies directed specifically against an epitope of the peptide.

In the second case, if the set of controls consisting of blood donors in apparent good health is 5 taken as a reference basis, without excluding the sera which are, on the face of it, seropositive, the standard deviation of the "non-MS controls" is 0.116. The threshold value then becomes  $0.13 + (3 \times 0.06) = 0.31$ .

According to this latter analysis, the test is 10 specific for MS. In this respect, it is seen that the test is specific for MS, since, as shown in Table 1, no control has a net OD above this threshold. In fact, this result reflects the fact that the antibody titres in patients suffering from MS are, for the most part, higher than in 15 healthy controls who have been in contact with MSRV-1.

In accordance with the first method of calculation, and as shown in Figure 41 and in Table 3, 6 of the 15 MS sera give a positive result (OD greater than or equal to 0.2), indicating the presence of IgGs 20 specifically directed against the S24Q peptide, hence against a portion of the reverse transcriptase enzyme of the MSRV-1 retrovirus encoded by its pol gene, and consequently against the MSRV-1 retrovirus.

Thus, approximately 40% of the MS patients 25 tested have reacted against an epitope carried by the S24Q peptide and possess circulating IgGs directed against the latter.

Two out of 15 blood donors in apparent good health show a positive result. Thus, it is apparent that 30 approximately 13% of the symptomless population may have been in contact with an epitope carried by the S24Q peptide under conditions which have led to an active immunization which manifests itself in the persistence of specific serum IgGs. These conditions are compatible with 35 an immunization against the MSRV-1 retrovirus reverse transcriptase during an infection with (and/or reactiva-

tion of) the MSRV-1 retrovirus. The absence of apparent neurological pathology recalling MS in these seropositive controls may indicate that they are healthy carriers and have eliminated an infectious virus after immunizing 5 themselves, or that they constitute an at-risk population of chronic carriers. In effect, epidemiological data showing that a pathogenic agent present in the environment of regions of high prevalence of MS may be the cause of this disease imply that a fraction of the population free 10 from MS has necessarily been in contact with such a pathogenic agent. It has been shown that the MSRV-1 retrovirus constitutes all or part of this "pathogenic agent" at the source of MS, and it is hence normal for controls taken from a healthy population to possess IgG 15 type antibodies against components of the MSRV-1 retrovirus.

Lastly, the detection of anti-S24Q antibodies in only one out of two MS cases tested here may reflect the fact that this peptide does not represent an 20 immunodominant MSRV-1 epitope, that inter-individual strain variations may induce an immunization against a divergent peptide motif in the same region, or that the course of the disease and the treatments followed may modulate over time the antibody response against the S24Q 25 peptide.

TABLE NO. 3

	CONTROLS	MS
5	0.101	0.136
	0.058	0.391
	0.126	0.37
	0.131	0.119
	0.105	0.267
	0.294	0.141
	0.116	0.102
	0.088	0.18
	0.1 05	0.411
10	0.172	0.164
	0.137	0.049
	0.223	0.644
	0.08	0.268
	0.073	0.065
	0.132	0.074
	Mean	0.129
	Std. Dev.	0.06
15	Threshold	0.31

d) Detection of anti-MSRV-1 IgM antibodies by  
ELISA:

20 The ELISA technique with the S24Q peptide was used to test for the presence of anti-MSRV-1 IgM specific antibodies in the same sera as above.

25 Figure 42 shows the results for each serum tested with an anti-IgM antibody. Each vertical bar represents the net optical density (OD at 492 nm) of a serum tested. The ordinate axis gives the net OD at the top of the vertical bars. The first 15 vertical bars lying to the left of the vertical line cutting the abscissa axis represent the sera of 15 healthy controls (blood donors), 30 and the vertical bars lying to the right of the vertical broken line represent the sera of 15 cases of MS tested.

The mean of the OD values for the MS cases tested is 1.6.

35 The mean of the net OD values for the controls is 0.7.

The standard deviation of the negative controls is 0.6.

The threshold of theoretical positivity may be calculated according to the formula:

5

threshold value = (mean of the OD values of the negative controls) + (3 x standard deviation of the OD values of the negative controls)

10 The threshold value is hence equal to  $0.7 + (3 \times 0.6) = 2.5$ ;

The negative results represent a non-specific "background" of the presence of antibodies directed specifically against an epitope of the peptide.

15 According to this analysis, and as shown in Figure 42 and in the corresponding Table 4, the IgM test is specific for MS, since no control has a net OD above the threshold. 6 of the 15 MS sera produce a positive IgM result

20 The difference in seroprevalence between the MS and control populations is extremely significant: "chi-squared" test,  $p < 0.002$ .

These results point to an aetiopathogenic role of MSRV-1 in MS.

25 Thus, the detection of IgM and IgG antibodies against the S24Q peptide makes it possible to evaluate, alone or in combination with other MSRV-1 peptides, the course of an MSRV-1 infection and/or of the viral reactivation of MSRV-1.

TABLE NO. 4

	CONTROLS	MS
5	1.449	0.974
	0.371	6.117
	0.448	2.883
	0.456	1.945
	0.885	1.787
	2.235	0.273
	0.301	1.766
	0.138	0.668
	0.16	2.603
	1.073	0.802
10	1.366	0.245
	0.283	0.147
	0.262	2.441
	0.585	0.287
	0.356	0.589
15	Mean	0.7
	Std. Dev.	0.6
	Threshold	
	Value	2.5

It is possible, as a result of the new discoveries made and the new methods developed by the inventors, to permit the improved implementation of diagnostic tests for MSRV-1 infection and/or reactivation and to evaluate a therapy in MS and/or RA on the basis of its efficacy in "negativizing" the detection of these agents in the patient's biological fluids. Furthermore, early detection in individuals not yet displaying neurological signs of MS or rheumatological signs of RA could make it possible to institute a treatment which would be all the more effective with respect to the subsequent clinical course for the fact that it would precede the lesion stage which corresponds to the onset of the clinical disorders.

Now, at the present time, a diagnosis of MS or RA cannot be established before a symptomatology of lesions has set in, and hence no treatment is instituted before the emergence of a clinical picture suggestive of lesions which are already significant. The diagnosis of an MSRV-1 and/or MSRV-2 infection and/or reactivation in man is

hence of decisive importance, and the present invention provides the means of doing this.

It is thus possible, apart from carrying out a diagnosis of MSRV-1 infection and/or reactivation, to 5 evaluate a therapy in MS on the basis of its efficacy in "negativizing" the detection of these agents in the patients' biological fluids.

**EXAMPLE 18 :**

10 1) MATERIALS AND METHODS

- Patients and clinical samples

Choroid plexus cells from MS patients and controls were obtained from the brain-cell library, Laboratoire R. Escourrolles, Hôpital de la Salpêtrière, 15 Paris, France. Non-tumoral leptomeningeal cells from controls were obtained as previously described (26). Peripheral blood from MS and control patients used for obtaining B-cell lines and plasma, were obtained from the Neurological Departments, CHU de Grenoble, and from 20 INSERM U 134, Hôpital de la Salpêtrière, France. Clinical details and origin of the 10 MS patients and of the 10 patients with other neurological diseases who provided CSF samples are given in Table 6.

- Cell cultures, virus isolation and purification

25 All cell-types were cultured as previously described (3, 5, 26).

All cultures were regularly screened for mycoplasma contamination with an ELISA mycoplasma-detection kit (Boehringer). No cell-extract nor supernatant used 30 contained detectable mycoplasma.

Extracellular virion purification and sucrose density gradients were performed as previously described (3, 5, 26). From each sucrose gradient 0.5-1ml fractions were collected from the top of the tubes, with a 1000 $\mu$ l 35 Pipetman and a different sterile tip for each fraction. 60 $\mu$ l were used for RT activity assay and the rest was

mixed with 1 volume of buffer containing 4M guanidinium thiocyanate, 0.5% N-Lauroyl sarcosin, 25mM EDTA, 0.2%  $\beta$ -mercaptoethanol adjusted at pH 5.5 with acetic acid. These mixtures were frozen at -80°C for further RNA extraction  
5 or directly processed according to Chomczynski (20), with an overnight precipitation step at -20°C, in presence of RNase-free glycogen (Boehringer). RNA was dissolved 20 to 50 $\mu$ l of DEPC-treated water in the presence of 1-2 $\mu$ l of recombinant RNase-inhibitor (Promega) and 0,1mM DTT. 10 $\mu$ l  
10 aliquots were used for each RT-PCR.

- Reverse transcriptase activity

RT-activity was tested with 20mM Mg<sup>++</sup> and poly-Cm or polyC templates, in virion pellets or fractions from sucrose gradients as previously described (3, 5, 26).

15 - cDNA synthesis and 'Pan-retro' RT-PCR with degenerate primers

A total RT-activity between 10<sup>6</sup>-10<sup>7</sup> dpm was required in the fraction containing the peak of purified virions. The "Pan-retro" RT-PCR technique (27) was  
20 performed on virion RNA extracted by the method of Chomczynski (20) and dissolved in 20  $\mu$ l RNase-free water.  
5  $\mu$ l RNA solution was incubated for 30 min at 37°C with 0.3 units (3 units for CSF series) of RNase-free DNase-1 (Boehringer) in a 20  $\mu$ l reaction containing 7.5 mM random  
25 hexamers, 5 mM Hepes-HCl pH 6.9, 75 mM KCl, 3 mM MgCl<sub>2</sub>, 10 mM DTT, 50 mM Tris-HCl pH 7.5, 0.5 mM each dNTP, and 20 units recombinant RNase inhibitor (Promega). The DNase was then heat inactivated at 80°C for 10 min. 20 units MoMLV RT (Pharmacia) and a further 20 units of RNase inhibitor  
30 were added to each tube in a Genesphere™ enclosure (Safetech, Ireland) and cDNA was synthesised for 90 min at 37°C. Following reverse transcription, the cDNA was boiled for 5 min then cooled rapidly on ice. The Round 1 PCR mix (final volume 25  $\mu$ l per reaction; 20 mM Tris-HCl pH 8.4,  
35 60 mM KCl, 2.5 mM MgCl<sub>2</sub>, 200 ng each of primers PAN-UO and PAN-DI [see Figure 44], 0.2 mM each dNTP) was treated with

0.3 units DNase-1 and then heat inactivated as above. 2.5  $\mu$ l cDNA was added in the Genesphere<sup>TM</sup> enclosure and the tubes heated to 80°C before adding 0.5 units Tag polymerase (Perkin Elmer) individually to each tube ("hot start"). Round 1 PCR parameters were 35 cycles of 95°C for 1 min, 34°C for 30 sec, 72°C for 1 min, with a final 7 min extension at 72°C. 0.5  $\mu$ l of Round 1 PCR product was transferred to the Round 2 DNase-treated PCR mix (composition as for Round 1 but containing primers PAN-UI and PAN-DI) using the "hot start" procedure. Round 2 PCR parameters were as for Round 1 but using 30 cycles only and annealing at 45°C for 1 min.

- Cloning of PCR products

PCR products were cloned using the TA-cloning<sup>®</sup> kit (British Biotechnology) according to the manufacturer's recommendations.

- Sequencing

Sequencing reactions were performed using the "Prism ready reaction kit dye deoxyterminator cycle sequencing kit" (Applied Biosystems). Automatic sequence analysis was performed on an automatic sequencer (Applied Biosystems, 373 A).

- RT-PCR with ST1 primer sets

The first PCR round was performed directly from the cDNA reaction mixture according to the one-step RT-PCR technique described by Mallet et al. (28). This one-step RT-PCR procedure reduced the probability of airborne contamination when opening the tubes and transferring PCR reagents after an independent cDNA synthesis. RNA was extracted as previously from 2ml of plasma (snap-frozen in liquid nitrogen and stored at -80°C) or from a 500  $\mu$ l sucrose fraction with a total RT-activity above 10<sup>6</sup> dpm, and resuspended in 50  $\mu$ l of RNase-free water. For each RT-PCR reaction 10  $\mu$ l of RNA solution was incubated in a Perkin-Elmer 480 thermocycler, 15 min at 20°C with 1U of RNase-free DNASE 1 and 1.2  $\mu$ l of 10X DNASE buffer (50mM

100

Tris, 10mM MgCl<sub>2</sub> and 0,1mM DTT) containing 1U/μl of RNase-inhibitor (PROMEGA), and heated at 70°C for 10 min for DNase inactivation. The solution was placed on ice and mixed (in conditions preventing airborne dust/DNA contamination) with 88 μl of PCR mix containing: 1X taq buffer, 25 nM/tube dNTPs, 40pM/tube of each first round primer (ST1.1 upstream primer: 5' AGGAGTAAGGAAACCCAACGGAC 3' (SEQ ID NO:99); ST1.1 downstream primer: 5' TAAGAGTTGCACAAGTGCG 3' (SEQ ID NO:100)), 2.5U/tube of taq (Appligene) and 10U/tube of AMV-RT (Boehringer). Each tube was further incubated in a Perkin-Elmer 480 thermocycler for 10 min at 65°C, followed by 2h at 42°C for cDNA synthesis and 5 min at 95°C for inactivation of AMV-RT and DNA denaturation. First round parameters were 40 cycles of 95°C for 1 min, 53°C for 2.5 min, 72°C for 1 min, with a final extension of 10 min at 72°C. 10μl of the first round were transferred to the second round PCR mix previously treated at 20°C for 15 min with RNase-free DNase 1 (0.02U/μl) followed by DNase inactivation at 70°C for 10 min. This mix contained 1X taq buffer, 25 nM/tube dNTPs, 40pM/tube of each second round primers [ST1.2 upstream primer: 5' TCAGGGATAGCCCCCATCTAT 3' (SEQ ID NO:101); ST1.2 downstream primer: 5' AACCCCTTGCCACTACATCAATT 3' (SEQ ID NO:102)] and 2.5U/tube of taq (Appligene). Second round parameters were 30 cycles of 95°C for 1 min, 53°C for 1.5 min, 72°C for 1 min, with a final extension of 8 min at 72°C. 20μl of this nested RT-PCR product were deposited on a 0,7% agarose gel containing ethidium bromide and exposed to UV light for the visualization of amplified products.

- Hybridisation analysis of PCR products: MSRV-pol detection by ELOSA

The protocol was essentially as previously described (21) but with the following modifications: Nunc Maxisorb microtitre plates were coated with 100 ng per well capture probe CpV1b (see Figure 44) either by passive

adsorption (21) or alternatively by using streptavidin coated plates and biotinylated CpV1b. Peroxidase-labelled detector probe DpV1 (see Figure 44) was used and the assay cut-off was defined as the mean of 4 negative controls 5 plus 0.2 OD<sub>492</sub> units.

- RNA extraction, cDNA synthesis and PCR amplification from MS plasma samples :

Total RNA was extracted from human MS plasma by a guanidium method as described elsewhere (29). Total RNA 10 extracted from 100 ul of plasma, were treated with RNase-free DNase I (0.1U/μl; Boehringer Manheim, France) and reverse transcribed under the conditions recommended by the manufacturer, using Superscript reverse transcriptase (Gibco-BRL, FRANCE). The resulting cDNAs were amplified by 15 semi-nested PCR through 35 cycles (94°C 1 min, 55°C 1 mn, 72°C 1 min 30 sec) and 72°C 8 min for a final extension.

Three different fragments in the RT region were amplified by the following specific primers :

- in the protease (PRT) region, for the 1st and 20 2nd round of PCR, respectively, sense primer [5' TCC AGC AGC AGG ACT GAG GGT 3' (SEQ ID NO:103)] and antisense primers [5' CTG TCC GTT GGG TTT CCT TAC TCC T 3' (SEQ ID NO:104) / 5' GAC AGC AAA TGG GTA TTC CTT TCC 3' (SEQ ID NO:105)]

25 - in the fragment A of the RT region (Cf. Fig 46), for the 1st and 2nd round of PCR, respectively, sense primer [5' AGG AGT AAG GAA ACC CAA CGG ACA G 3' (SEQ ID NO:106)] and antisense primers [5' TGT ATA TAA TGG TCT GGC TAT TGG G 3' (SEQ ID NO:107) / 5' TTC GGC AGA AAC CTG TTA 30 TGC CAA GG 3' (SEQ ID NO:108)]

- in the fragment B of the RT region (Cf. Fig. 46), for the 1st and 2nd round of PCR, respectively, sense primers [5' GGC TCT GCT CAC AGG AGA TTA GAT AC 3' (SEQ ID NO:109) / 5' AAA GGC ACC AGG GCC CTC AGT GAG GA 3' (SEQ ID 35 NO:110)] and antisense primer 3'[5' GGT TTA AGA GTT GCA CAA GTG CGC AGT C 3' (SEQ ID NO:101)].

The amplified fragments were analysed on ethidium bromide-stained agarose gels, cloned in TA cloning vector (Invitrogen) and sequenced.

## 2) RESULTS

5 - Specific retroviral RNA is found in extracellular virions from MS patient-derived cell cultures and in MS patients' CSF.

Choroid plexus cells (4) (obtained post-mortem) and EBV-immortalized peripheral blood B-lymphocytes (30, 10 31) from MS patients gave rise to cultures expressing 100-120 nm viral particles associated with RT-activity similar to that of the original LM7 isolate (3). Similar cell-types from non-MS donors produced neither this RT-activity nor virions. All the 'infected' cultures were poorly 15 and/or transiently productive and/or had a limited lifespan. Therefore, in order to analyse the genomic RNA present in the very limited quantity of extracellular virions, we used an RT-PCR approach to amplify, with degenerate primers, a conserved region of the pol gene 20 present in all known retroviruses (12); the techniques based on this approach will be called "Pan-retro" RT-PCR. Extensive DNase treatment of samples and reagents was essential, because human DNA contains many endogenous retroviral elements amplifiable by this technique.

25 "Pan-retro" RT-PCR experiments were performed on sucrose-density gradient purified virions from supernatants of different types of cell cultures and their non-infected controls: (i) choroid plexus cells sampled post-mortem from MS brain (PLI-1), (ii) choroid plexus cells from non-30 MS brain autopsy, infected by co-culture with irradiated LM7 cells (LM7P), and (iii) identical non-infected choroid-plexus cells. "Early" B-cell lines obtained by spontaneous in vitro transformation of two EBV-seropositive individuals, (iv) one MS patient and (v) one 35 non-MS control, were also analysed. Figure 43 illustrates the RT-activity in sucrose-gradient fractions obtained

from the B-cell cultures. The technique described by Shih et al. (12) was modified in a semi-nested RT-PCR protocol (27) using degenerate primers (Fig.2) and extensive DNase treatment. PCR amplifications were performed in London 5 (Dpt of Virology, U.C.L.M.S.) on coded aliquots of the density gradient fractions. Blind and systematic cloning and sequencing of the PCR products were undertaken in an independent laboratory (bioMérieux, Lyon). After complete sequencing of 20 to 30 clones per sucrose gradient 10 fraction, the codes were broken and results analysed in parallel with the RT-activity data.

Table 5 presents the distribution of sequences obtained from sucrose gradient fractions containing the peak of viral RT-activity in MS-derived cultures and also the 15 sequences amplified from the corresponding RT-activity negative fractions of uninfected cultures. The predominant sequence detected in bands of the expected size ( $\sim$ 140 bp) amplified in all the RT-activity positive fractions (but not in the RT-activity negative fractions) was different 20 from known retroviruses and was designated MSRV-cpol. MSRV-cpol sequences exhibited partial homology (70-75%) with ERV9, a previously described endogenous retroviral sequence (18). A few ERV9 sequences (>90% homology with ERV9) were also present but clearly represented a minority 25 of clones. In addition to typical pol sequences, numerous PCR artefacts (primer multimers, concatemers or single-primer amplifications) related to the use of degenerate primers and low-temperature annealing, were found in all samples (Table 5).

30 Figure 44 shows an alignment of a consensus sequence of MSRV-cpol with the corresponding VLPQG / YMDD region of diverse retroviruses. Figure 45 displays a phylogenetic tree based on the evolutionarily conserved amino acid sequences of both exogenous and endogenous retroviruses in this 35 region. From this tree it can be seen that the pol gene of

MSRV is phylogenically related to the C-type group of oncovirinae.

A small scale study was performed to determine the prevalence of MSRV c-pol sequences in the CSF of patients 5 with MS. Identification of MSRV-cpol in PCR products by cloning and sequencing is both laborious and time consuming. We therefore devised an enzyme-linked oligosorbent assay (ELOSA), using a capture probe (CpV1B) and a peroxidase-labelled detector probe (DpV1), for the 10 rapid identification of MSRV-cpol sequences in 'Pan-retrovirus' PCR products (Figure 44). The specificity of this sandwich hybridisation-based assay for HMSRV-cpol was tested with both distantly related (HIV and MoMLV) and closely related (ERV9) pol sequences. No significant cross 15 reactivity with such targets was observed despite the ability of the ELOSA to detect as little as 0.01 ng of MSRV-cpol DNA.

Cerebrospinal fluid (CSF) samples were available from 10 patients with MS and from 10 patients with other 20 neurological disorders. Total RNA was extracted from CSF pellets, reverse transcribed and amplified as above. ELOSA analysis (Table 6) of the PCR products revealed MSRV-cpol sequences in 5 of the 10 MS patient samples but in none of the 10 samples from patients with other neurological 25 diseases ( $P<0.05$ ). The presence of MSRV-cpol did not appear to be correlated with age, sex or type of MS, but was seen in untreated patients only (5/6). No patient with immunosuppressive therapy was found positive (0/4). No correlation between MSRV-cpol detection and CSF cell count 30 was observed.

- Cloning and sequencing a larger region of the pol gene

An independent identification of the MSRV genomic sequence was obtained by a non-PCR approach using RNA extracted from concentrated virions derived from 2,5 35 liters of LM7-infected sub-cultures of choroid plexus cells. A limited number of clones was obtained by direct

cloning of the cDNA, one of which (PSJ17) showed partial homology with ERV9 pol. Specific primers based on the MSRV-cpol region and on the PSJ17 clone, amplified a 740 bp fragment linking the two independent sequences in RNA extracted from purified virions. PSJ17 was localised on the 3' side of MSRV-cpol. Further sequence extension on the 5' side of MSRV-cpol and on the 3' side of PSJ17, was obtained using RT-PCR approaches on RNA from purified LM7-like virions produced in MS choroid plexus cultures (4).

In Figure 46, the nucleotide sequence corresponding to overlapping clones obtained by sequence extension in the pol gene is represented with the aminoacid translation corresponding to the putative open reading frames (ORFs) of the protease and of the reverse-transcriptase. The active site motifs of the protease (PRT) and of the reverse-transcriptase (RT) are underlined. In the C-terminal region of the RT sequence, the dispersed amino acid residues regularly present in retroviral RNase H domains, are also underlined.

- Non-degenerate primers detect MSRV-specific RNA in virions associated with the peak of RT-activity . and in in MS patients' plasma

PCR primers (ST1.1 primer set; positions 603-625/1732-1714, on Fig.4) based on overlapping clones in the pol gene, amplified a 1.15 kb segment of the RT region from several different isolates obtained from different MS patients. Nested primers (ST1.2; positions 869-889/1513-1490, on Fig.46) generated a 700 bp fragment (Figure 47) which was more easily visualised by ethidium bromide staining than the first round product generated by ST1.1. The specificity of PCR products was confirmed by stringent hybridisation with a peroxidase-labeled MSRV-cpol probe (Fig.44), using the ELOSA technique (21).

The ST1.1 and 2 primer set was used to detect extracellular MSRV RNA in human plasma, although non-optimal for this application. Figure 47 illustrates the

results of PCR amplification of cDNA derived from 2 MS patient and 2 control plasma samples tested in parallel with cDNA from the sucrose density gradient fractions of an MS choroid plexus isolate. Taq-sequencing of the 700 bp bands confirmed the presence of MSRV sequence. A very faint 700 bp band is also visible in fraction 10 which corresponds to the bottom of the tube where aggregated particles usually sediment. Control RT-PCR for cellular aldolase transcripts on plasma-derived RNA was negative, indicating that the results were not due to cellular RNA released by cell lysis during plasma separation. It should be noted that this PCR technique was not designed for epidemiological studies since its sensitivity is impaired by the length of the cDNA required (1.15 kb).

Non degenerate primers amplifying three fragments of the pol gene (the whole protease region, regions A and B of the reverse transcriptase; Cf. Fig. 46) were also used to confirm the presence of MSRV sequences in DNase-treated RNA from MS plasma. These fragments were amplified from the plasma of a further 4 MS patients with active disease. Sequence analysis confirmed that the PRT and RT regions were homologous (>95% and >90% respectively) to MSRV sequences previously obtained on culture virion. No such sequence were detected in plasma from healthy controls (n=4), tested in parallel with MS plasma.

### 3) DISCUSSION

#### - Phylogeny of MSRV

From the results of this study, it can be concluded that the virus previously referred to as "LM7" (3, 5, 26) posseses an RNA genome containing the MSRV pol sequences described here.

The conserved RT motif of both MSRV and ERV9 is two amino acids shorter than that of other retroviruses, apart from human foamy viruses which nonetheless have a functional RT. The potential ORF encompassing the entire PRT-RT

region is consistent with the virion-associated RT-activity detected in sucrose density gradients with infected culture supernatants. Moreover, since we have recently succeeded in expressing a recombinant protein 5 from the sequence of MSRV protease cloned from MS plasma, we can confirm the reality of the potential PRT ORF. Similar cloning and expression of other sequences containing potential ORFs for MSRV proteins, is being undertaken to confirm their ability to encode enzymes and 10 structural proteins of MSRV virions.

The phylogenetic tree in Figure 45, based on the most conserved amino acid sequence in retroviruses (VLPQG...YXDD), shows that the MSRV pol gene is related to the C-type oncoviruses. Apart from ERV9, the closest known 15 retroviral element is RTL-H, a human endogenous sequence known to have a subtype with a functional pol gene (32). In the pol region, this phylogenetic affiliation to C-type oncoviruses apparently contradicts our previous assumptions based on the general morphology of the 20 particles observed by electron microscopy (EM), which were compatible with a B or D-type oncovirus (3, 5, 26). However, preliminary data on env sequences detected in MSRV virions, would suggest a greater phylogenetic proximity 25 to D-type. Such difference in phylogenies of the pol and env genes have been described in MPMV and suggest a recombinatorial origin in D-type retroviruses (33). D to C type morphological conversion is also possible since it has been reported that a single amino acid substitution in the gag protein can convert retrovirus morphology to that 30 of a different type (34).

- Is MSRV an exogenous retrovirus sharing extensive homology with a related endogenous retrovirus family or an endogenous retrovirus producing extracellular virions?

Southern blot analysis with an MSRV pol probe 35 under stringent conditions, showed hybridisation with a multicopy endogenous family (data not presented),

indicating the existence of endogenous elements more closely related to MSRV than ERV9 itself. Consequently, we were unable to look for a virion-specific provirus in MSRV-producing cells. In agreement with southern blot 5 findings, PCR studies on genomic DNA showed multiple band amplification of MSRV-related endogenous sequences. Since pol is the most conserved retroviral gene, the sequence described here is the least suitable region to discriminate between exogenous and endogenous sequences. 10 It is hoped that sequence information from other parts of the genome may permit such a discrimination, would it be on a tiny portion as has recently been demonstrated for the Jaagsiekte retrovirus (JSRV) of sheep (35). With such sequence data, it would then become possible to identify 15 the MSRV-specific provirus in the genome of virion-producing cell cultures.

MSRV could represent a virion-producing exogenous member of an ERV9-like endogenous family, just as exogenous strains exist in the well-studied mouse mammary tumour 20 virus (MMTV) and murine leukaemia virus (MuLV) retroviral families of mice, and also, in the JSRV retroviral family of sheep (36). Alternatively, it is also conceivable that the extracellular MSRV virions may be produced by a replication-competent endogenous provirus. Whether MSRV is 25 exogenous or endogenous, conceptual similarities exist with the category of retroviruses represented by MuLV, MMTV and JSRV. Unlike defective endogenous elements, this category of agents are known to produce infectious and pathogenic virions, to cause neurological disease (37), 30 solid tumours / leukaemias (36, 38) and to express "endogenous superantigens" (39, 40). Furthermore, in MuLV infections, the genetic endogenous retroviral background of the mouse strain can determine susceptibility or resistance to disease (39, 41). Indeed, such interactions 35 between an infectious retrovirus and its endogenous counterpart may be relevant in the pathogenesis of MS,

since endogenous retroviral genotypes are not identical in all individuals. A genetic control due to related endogenous retroviral genotypes could therefore contribute to the known hereditary susceptibility to MS (43), if MSRV 5 does indeed play an active role in this disease.

Elsewhere, the data in Table 5 suggest that ERV9 elements may be co-expressed, possibly via trans-activation in infected cells, and give rise to heterologous RNA packaging in MSRV virions. Such heterologous packaging is 10 known to occur in other retroviral systems (42).

- A role for the numerous common viruses previously evoked in MS ?

Among the numerous reports of viruses putatively involved in the aetiopathogenesis of MS, a significant 15 proportion focus on two viral families, the paramyxoviridae and the herpesviridae. Regarding the paramyxoviridae, the key observation is of a frequently increased antibody titer to measles virus in MS patients essentially directed, in CSF, against measles fusion 20 protein (44). The existence of aminoacid similarities between conserved domains of the fusion proteins of paramyxoviridae and the transmembrane protein of retroviruses (45), may explain this observation if antigenic cross-reactivity between these two proteins 25 occurred.

With regard to the herpesvirus family, the involvement of Epstein-Barr Virus (EBV), Herpes Simplex Virus type 1 (HSV-1) and, most recently, Human Herpes Virus 6 (HHV-6) has been proposed (31, 46, 47). From our previous studies 30 and from those of other groups, it appears that herpesviruses may play an important role in MSRV expression: we have shown that HSV-1 immediate-early ICP0 and ICP4 proteins can transactivate MSRV/LM7 in vitro (6) and Haahr et al. have proposed an important 35 epidemiological role for EBV, as a co-factor in MS, triggering retrovirus reactivation (31). The recent

description by Challoner et al. (47) showing significant expression of HHV6 proteins in MS plaques may also suggest a similar role for HHV6 in the brain.

5 EXAMPLE 19 : MSRV GENOME DETECTION TECHNIQUE

Following 0.4 µm filtration to remove cellular debris and RNase digestion to remove residual non-encapsidated RNA, serum was processed to extract viral RNA by means of adsorption to a silica matrix. Viral RNA was 10 subjected to DNase digestion, then a combined reverse transcription-PCR (RT-PCR) reaction was performed using primers PTpol-A (sense: 5'xxxx3', SEQ ID NO:183) and PTpol-F (antisense: 5'xxxx3', SEQ ID NO:184). A second round of amplification with nested primers PTpol-B (sense: 15 5'xxxx3', SEQ ID NO:185) and PTpol-E (antisense: 5'xxxx3', SEQ ID NO:186) generated a 435 bp PCR product which was identified by gel electrophoresis. The specificity of each product was confirmed by dideoxy sequencing. Control reactions without reverse transcriptase were performed to 20 ensure that the products were derived from viral RNA. In addition, to exclude the possibility that the extracted viral RNA might be contaminated with host cell derived nucleic acids, aliquots were tested by nested PCR for the presence of pyruvate dehydrogenase (PDH) DNA and RNA. 25 Samples which generated a signal in either the PDH or the "no-RT" PCR assays were excluded from the analysis.

Sera from patients with clinically active MS and controls were amplified by RT-PCR and sequenced. Virion associated MSRV-RNA was detected in the serum of 10 of 19 30 (53%) patients with MS but in only 3 of 44 controls without MS ( $P=0.0001$ ). The control group consisted of 8 patients (all MSRV-RNA negative) with rheumatological disorders and 36 healthy adults. MSRV-RNA titres in both MS patients and controls were apparently low because even 35 moderate dilution of sera (<10 fold) caused loss of signal.

In MS patients, detection of MSRV-RNA was not associated with age, sex, disease duration, or MS type, however a significant negative correlation with treatment was observed. 26 serum samples were obtained from the 19 5 patients ; 100% of the sera from untreated patients contained detectable MSRV-RNA whereas it was detectable in only 4 of 19 samples (21%) obtained during treatment with corticosteroids and/or azathioprine ( $P=0.001$ ).

The reason for the apparent loss of virion 10 associated MSRV-RNA during immunosuppressive treatment is unknown but the finding is in agreement with the previous observations on the detection of MSRV in cerebrospinal fluid.

15

TABLE 7  
DETECTION OF VIRION ASSOCIATED MSRV-RNA IN MS UNTREATED PATIENTS & CONTROLS

	Positive	Negative	Total	% Positive
Controls without MS <sup>a</sup>	3 <sup>b</sup>	41	44	7%
MS sera untreated at time of sampling	7	0	7	100%

20 a The control group consisted of 8 patients with miscellaneous non-MS disorders and 36 healthy adults.

b The detection of MSRV RNA in plasma of a few controls in conditions which select virion-packaged RNA, is consistent with the knowledge that a virus associated with MS should 25 be present in a minor proportion of apparently healthy population. Indeed, such individuals can be either healthy carriers or be in the pre-clinical (or sub-clinical) phase of the disease which can last for years.

## METHOD :

- Modified SNAP RNA extraction with filtration and RNase digestion

(All centrifugations are at room temperature)

5 Up to 500 microlitres of serum is filtered using 0.45 micron spin filters (Nanosep MF from Flowgen Catalogue No. U3-0126 Ref. ODM45). The serum is spun for 5 min at 130,000 g (or for further 10 min if necessary).

10 150 microlitres of filtered serum is incubated with 10 units RNase One (Promega Catalogue No.M4261) for 30 min at 37°C.

The 150 microlitres was then extracted using the SNAP RNA extraction kit (Invitrogen) as below:

15 - 10 micrograms of poly A RNA was added to the 450 microlitres of Binding Buffer to act as a carrier ; this was then added to the serum and mixed by inversion 6 times ; 300 microlitres of propan-2-ol was then added and mixed by inversion 10 times ; 500 microlitres was transferred to the SNAP column and spun at 1300 g for 20 1 min and the flow-through discarded ; the remainder was then added to the SNAP column and spun at 1300 g for 1 min and the flow-through discarded ; the column was then washed with 600 microlitres of Super wash and the flow-through discarded ; the column was then washed with 600 25 microlitres of 1x RNA wash and the flow-through discarded ; this wash was repeated with a 2 min 1300 g spin and the flow-through discarded ; the bound nucleic acid was then eluted by incubating with 135 microlitres of RNase free water for 5 min and spun at 1300 g for 1 min.

30 - 15 microlitres of 10x DNase buffer and 3 microlitres (30 units) of DNase I, RNase free (Boehringer Mannheim Cat. No. 776 785) was added and incubated for 30 min at 37°C ; 450 microlitres of Binding Buffer was added and mixed by inversion 6 times ; 300 microlitres of 35 propan-2-ol was then added and mixed by inversion 10 times ; 500 microlitres was transferred to the SNAP column

and spun at 1300 g for 1 min and the flow-through discarded ; the remainder was then added to the SNAP column and spun at 1300 g for 1 min and the flow-through discarded ; the column was then washed with 600 5 microlitres 1x RNA wash and the flow-through discarded ; this wash was repeated with a 2 min 1300 g spin and the flow-through discarded ; the bound nucleic acid was then eluted by incubating with 105 microlitres of RNase free water for 5 min and spun at 1300 g for 1 min.

10

- Titan RT-PCR

RT-PCR was performed using the Titan one tube RT-PCR system (Boehringer Mannheim Cat. No. 1 855 476) 25 microlitres of RNA was used in the combined RT-PCR 15 reaction. The total reaction volume was 50 microlitres. Promega rRNAsin (10 units) was the RNase inhibitor used. 170 ng of primers SEQ ID NO:183 and SEQ ID NO:184, respectively, were used. A single master mix was prepared and the sample RNA added last. This was performed at room 20 temperature, not on ice.

The RT step consisted of two sequential 30 min incubations at 50°C and then 60°C. This was immediately followed by the PCR which had the following steps.

\* Initial denaturation of template at 94°C for 2 min,  
25 \* 40 cycles of 94°C for 30 seconds ; 60°C for 30 seconds ;  
68°C for 45 seconds,  
\* 1 cycle of 68°C for 7 min.

The second round PCR was performed using the Expand long template PCR system (Boehringer Mannheim Cat. 30 No. 1681 842). 0.5 microlitres of the RT-PCR mix was added to 25 microlitres of the round 2 PCR mix. Buffer No. 3 and 50 ng of primers B and E were used. The PCR had the following steps:

\* 5 cycles of 94°C for 30 seconds, 60°C for 30 seconds.,  
35 68°C for 45 seconds,  
\* 1 cycle of 68°C for 7 min.

The PCR products were then run on a 2% agarose gel.

The no RT controls were performed using "Expand" PCR system for both rounds. The first round was 40 cycles  
5 and the second round 20 cycles.

As a positive control a DNA dilution series was used in both the RT-PCR and the "no RT" PCR. For a result to be valid the RT-PCR and "no-RT" PCRs had to have detected DNA equivalent to between 1 and 0.1 cells.

10 The analysis of PCR products of an approximately 435 bp fragment in the pol region is shown in Table 8.

TABLE 8  
ANALYSIS OF PCR PRODUCTS WITH ORF \*

15

Exp	Disease	Clone	ORF	Fragment (bp)	AA-RT Motif Site
46-7	MS	1	+	429	YGDD
		5	+	429	YGDD
		8	+	429	YGDD
20	68-1	41	+	438	YMDD
		42	+	438	YMDD
		43	+	438	YMDD

25 \* Defective RNA can also be present in circulating virions, since the fidelity of the MSRV reverse transcriptase appears to be low and since recombination events with related endogenous elements can occur. It is then obvious that the intra- and inter- patients  
30 variability can be greater than that illustrated in this example, because of these encapsidated defective MSRV RNA copies.

Table 9 which data have been determined from the  
35 alignments of Figures 49 to 53, shows a variability :

- between the clones obtained from the same patient plasma sample in the same PCR amplification experiment ; this means that the patient possesses a virion population which comprises different MSRV variants at a given time,
- 5 - between the sequenced variant populations from different patients ; this means that the variants differ from a patient to another patient.

TABLE 9

10      Degree of identity (percentage) between nucleotide sequences and between peptide sequences,  
by direct comparison of said sequences (see Figures 49-53)

Patient	68-1	46-7
Nucleotide sequences	between SEQ ID NO:169 and MSRV-pol (SEQ ID NO:1) 90,4 % b 92,3 % a  SEQ ID NOS:170, 171, 172 between them 98,6 % b 98,7 % a	between SEQ ID NO:176 and MSRV-pol (SEQ ID NO:1) 82,5 % a 84 % b  SEQ ID NOS:177, 178, 179 between them 94,5 % a 95,1 % b
Peptide sequences	between SEQ ID NOS:173, 174, 175 and SEQ ID NO: 81 %  SEQ ID NOS:173, 174, 175 between them 97 %	between SEQ ID NOS:180, 181, 182 and SEQ ID NO: 73,5 %  SEQ ID NOS:180, 181, 182 between them 89 %

- 15 a) this percentage is determined on the basis of sequences excluding the primers  
 b) this percentage is determined on the basis of sequences including the primers.
- 20 From Figures 53A and 53B, the variability between tested patients sequences can be determined :

116

- between SEQ ID NO:169 and SEQ ID NO:176 : 16,5 %<sup>a</sup> and 14,8 %<sup>b</sup>
- between the peptide sequences obtained from SEQ ID NO:169 and SEQ ID NO:176 : 20 %.

5

Four microorganisms are mentioned in the specification page 3 lines 15-26 and they are identified below. They have all been deposited with the ECACC\*, in accordance with the provisions of the Budapest Treaty.

10

- LM7PC deposited on 22nd July 1992 under No. 92072201,
- PLI-2 deposited on 8th January 1993 under No. 93010817,
- POL-2 deposited on 22nd July 1992 under No. V92072202, and
- 15 - MS7PG deposited on 8th January 1993 under No. V93010816.

\* ECACC : European Collection of Animal Cell Cultures  
Vaccine Research and Production Laboratory

20

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25

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121

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## SEQUENCE LISTING

## (1) GENERAL INFORMATION:

5 (i) APPLICANT: BIO MERIEUX

10 (ii) TITLE OF THE INVENTION: VIRAL MATERIAL AND NUCLEOTIDE  
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PROPHYLACTIC AND THERAPEUTIC PURPOSES

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(A) MEDIUM TYPE: Floppy disk

(B) COMPUTER: IBM PC compatible

25 (C) OPERATING SYSTEM: PC-DOS/MS-DOS

(D) SOFTWARE: PatentIn Release #1.0, Version #1.30

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30 (B) FILING DATE:

(viii) ATTORNEY/AGENT INFORMATION:

(A) NAME: Dominique GUERRE

(B) REGISTRATION NUMBER:

35 (C) REFERENCE/DOCKET NUMBER: MD/B05B2679

123

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- (A) TELEPHONE: 4 72 69 84 30  
(B) TELEFAX: 4 72 69 84 31

## 5 (2) INFORMATION FOR SEQ ID NO: 1:

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 1158 base pairs  
(B) TYPE: nucleotide  
10 (C) STRANDEDNESS: single  
(D) TOPOLOGY: linear

## (ii) MOLECULE TYPE: cDNA

## 15 (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 1:

CCCTTGC	CA CTACATCAAT TTTAGGAGTA AGGAAACCCA ACGGACAGTG GAGGTTAGTG	60
CAAGAACTCA	GGATTATCAA TGAGGCTGTT GTTCCTCTAT ACCCAGCTGT ACCTAACCC	120
TATACAGTGC	TTTCCCAAAT ACCAGAGGAA GCAGAGTGGT TTACAGTCCT GGACCTTAAG	180
20 GATGCCTTT	TCTGCATCCC TGTACGTCC CACTCTCAAT TCTTGTTCGC CTTTGAAGAT	240
CCTTGAA	CC ACCGTCTCA ACTCACCTGG ACTGTTTAC CCCAAGGGTT CAGGGATAGC	300
CCCCATCTAT	TTGCCAGGC ATTAGCCAA GACTTGAGTC AATTCTCATCA CCTGGACACT	360
CTTGTCTTC	AGTACATGGA TGATTTACTT TAGTCGCC CTTCAGAAAC CTTGTGCCAT	420
CAAGCCACCC	AAGAACTCTT AACTTCCCTC ACTACCTGTG GCTACAAGGT TTCCAAACCA	480
25 AAGGCTCGGC	TCTGCTCACA GGAGATTAGA TACTNAGGGC TAAAATTATC CAAAGGCACC	540
AGGGCCCTCA	GTGAGGAACG TATCCAGCCT ATACTGGCTT ATCCTCATCC CAAAACCTA	600
AAGCAACTAA	GAGGGTCCCT TGGCATAACA GGTTCTGCC GAAAACAGAT TCCCAGGTAC	660
ASCCAATAG	CCAGACCATT ATATACACTA ATTANGAAA CTCAGAAAGC CAATACCTAT	720
TTAGTAAGAT	GGACACCTAC AGAAGTGGCT TTCCAGGCC TAAAGAAGGC CCTAACCCAA	780
30 GCCCCAGTGT	TCAGCTGCC AACAGGGCAA GATTTTCTT TATATGCCAC AGAAAAAACAA	840
GGAATAGCTC	TAGGAGTCCT TACCGCAGGTC TCAGGGATGA GCTTGCAACC CGTGGTATAC	900
CTGAGTAAGG	AAATTGATGT AGTGGCAAAG GGTTGCCCTC ATNGTTATG GTAAATGGNG	960
GCAGTAGCAG	TCTNAGTATC TGAAGCAGTT AAAATAATAC AGGGAAGAGA TCTTNCTGTG	1020
TGGACATCTC	ATGATGTGAA CGGCATACTC ACTGCTAAAG GAGACTTGTG GTTGTCA	1080
35 AACCATTTAC	TTAANTATCA GGCTCTATTA CTTGAAGAGC CAGTGCTGNG ACTGCCACT	1140
TGTGCAACTC	TTAAACCC	1158

## (2) INFORMATION FOR SEQ ID NO: 2:

## (i) SEQUENCE CHARACTERISTICS:

- 5                   (A) LENGTH: 297 base pairs  
                 (B) TYPE: nucleotide  
                 (C) STRANDEDNESS: single  
                 (D) TOPOLOGY: linear

10               (ii) MOLECULE TYPE: cDNA

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 2:

CCCTTGCCA	CTACATCAAT	TTTAGGAGTA	AGGAAACCCA	ACGGACAGTG	GAGGTTAGTG	60
15 CAAGAACTCA	GGATTATCAA	TGAGGCTGTT	GTTCCTCTAT	ACCCAGCTGT	ACCTAACCCCT	120
TATACAGTGC	TTTCCCAAAT	ACCAAGGAA	GCAGAGTGGT	TTACAGTCCT	GGACCTTAAG	180
GATGCCTTT	TCTGCATCCC	TGTACGTCC	GACTCTCAAT	TCTTGTTCGC	CTTGAAAGAT	240
CCTTGAAACC	CAACGTCTCA	ACTCACCTGG	ACTGTTTAC	CCCAAGGGTT	CAAGGGA	297

20

## (2) INFORMATION FOR SEQ ID NO: 3:

## (i) SEQUENCE CHARACTERISTICS:

- 25                   (A) LENGTH: 85 base pairs  
                 (B) TYPE: nucleotide  
                 (C) STRANDEDNESS: single  
                 (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

30               (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 3:

GTTCAGGGAT	ANCCCTCATC	TCTTGTC	GGTACTGGCC	CAAGATCTAG	GCCACTTCTC	60
AGGTCCAGSN	ACTCTGTYCC	TTCAG				85

## (2) INFORMATION FOR SEQ ID NO: 4:

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 86 base pairs
- 5 (B) TYPE: nucleotide
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

## (ii) MOLECULE TYPE: cDNA

10

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 4:

GTTCAGGGAT AGCCCCATC TATTGGCCA GGCCTAGCT CAATACTTGA GCCAGTTCTC	60
ATACCTGGAC AYTCTYGTCC TTCTGGT	86

15

## (2) INFORMATION FOR SEQ ID NO: 5:

## (i) SEQUENCE CHARACTERISTICS:

- 20 (A) LENGTH: 85 base pairs
- (B) TYPE: nucleotide
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

25 (ii) MOLECULE TYPE: cDNA

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 5:

GTTCARRGAT AGCCCCATC TATTGGCCW RGYATTAGCC CAAGACTTGA GYCAATTCTC	60
30 ATACCTGGAC ACTCTTGTC TTGTYRG	85

## (2) INFORMATION FOR SEQ ID NO: 6:

35 (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 85 base pairs

126

- (B) TYPE: nucleotide
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

5 (ii) MOLECULE TYPE: cDNA

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 6:

GTTCAGGGAT AGCTCCCATC TATTGCGCTT GGCATTAACC CGAGACTTAA GCCAGTTCTY	60
10 ATACGTGGAC ACTCTTGTCC TTTGG	85

(2) INFORMATION FOR SEQ ID NO: 7:

15 (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 111 base pairs
- (B) TYPE: nucleotide
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

20

(ii) MOLECULE TYPE: cDNA

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 7:

25

GTGTTGCCAC AGGGGTTTAR RGATANCYCY CATCTMTTG GYCWRGYAYT RRCYCRAKAY	60
YTRRGYCAVT TCTYAKRYSY RGSNAYTCTB KYCCTTYRGT ACATGGATGA C	111

30 (2) INFORMATION FOR SEQ ID NO: 8:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 645 base pairs
- (B) TYPE: nucleotide
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

35

(ii) MOLECULE TYPE: cDNA

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 8:

5

TCAGGGATAG	CCCCCATCTA	TTTGGCCAGG	CATTAGCCCA	AGACTTGAGT	CAATTCTCAT	60
ACCTGGACAC	TCTTGTCCCT	CAGTACATGG	ATGATTTACT	TTTAGTCGCC	CGTTCAGAAA	120
CCTTGTGCCA	TCAAGCCACC	CAAGAACTCT	TAACCTTCCT	CACTACCTGT	GGCTACAAGG	180
TTTCCAAACC	AAAGGCTCGG	CTCTGCTCAC	AGGAGATTAG	ATACTNAGGG	CTAAAATTAT	240
10 CCAAAGGCAC	CAGGGCCCTC	AGTGAGGAAC	GTATCCAGCC	TATACTGGCT	TATCCTCATC	300
CCAAAACCCCT	AAAGCAACTA	AGAGGGTTCC	TTGGCATAAC	AGGTTTCTGC	CGAAAACAGA	360
TTCCCAGGTA	CASCCAATA	GCCAGACCAT	TATATACT	AATTANGGAA	ACTCAGAAAG	420
CCAATACCTA	TTTAGTAAGA	TGGACACCTA	CAGAAGTGGC	TTTCCAGGCC	CTAAAGAAGG	480
CCCTAACCCA	AGCCCCAGTG	TTCAGCTTGC	CAACAGGGCA	AGATTTTCT	TTATATGCCA	540
15 CAGAAAAAAC	AGGAATAGCT	CTAGGAGTCC	TTACGCAGGT	CTCAGGGATG	AGCTTGCAAC	600
CCGTGGTATA	CCTGAGTAAG	GAAATTGATG	TAGTGGCAAA	GGGTT		645

(2) INFORMATION FOR SEQ ID NO: 9:

20

(i) SEQUENCE CHARACTERISTICS:

25

- (A) LENGTH: 741 base pairs
- (B) TYPE: nucleotide
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 9:

30

CAAGCCACCC	AAGAACTCTT	AAATTCCTC	ACTACCTGTG	GCTACAAGGT	TTCCAAACCA	60
AAGGCTCAGC	TCTGCTCACA	GGAGATTAGA	TACTTAGGGT	TAAAATTATC	CAAAGGCACC	120
AGGGGCTCA	GTGAGGAACG	TATCCAGCCT	ATACTGGGTT	ATCCTCATCC	CAAACCCCTA	180
AAGCAACTAA	GAGGGTTCCCT	TAGCATGATC	AGGTTTCTGC	CGAAAACAAG	ATTCCCAGGT	240
35 ACAACCAAAA	TAGCCAGACC	ATTATATACA	CTAATTAAGG	AAACTCAGAA	AGCCAATACC	300
TATTTAGTAA	GATGGACACC	TAAACAGAAG	GCTTCCAGG	CCCTAAAGAA	GGCCCTAACCC	360

128

CAAGCCCCAG TGTCAGCTT GCCAACAGGG CAAGATTTT CTTTATATGG CACAGAAAAA 420  
ACAGGAATCG CTCTAGGAGT CCTTACACAG GTCCGAGGGA TGAGCTTGCA ACCCGTGGCA 480  
TACCTGAATA AGGAAATTGA TGTAGTGGCA AAGGGTTGGC CTCATNGTTT ATGGGTAATG 540  
GNGGCAGTAG CAGTCTNAGT ATCTGAAGCA GTTAAAATAA TACAGGGAAG AGATCTTNCT 600  
5 GTGTGGACAT CTCATGATGT GAACGGCATA CTCACTGCTA AAGGAGACTT GTGGTTGTCA 660  
GACAACCATT TACTTAANTA TCAGGCTCTA TTACTTGAAG AGCCAGTGCT GNGACTGCGC 720  
ACTTGTGCAA CTCTTAAACC C 741

10 (2) INFORMATION FOR SEQ ID NO: 10:

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 93 base pairs
- (B) TYPE: nucleotide
- 15 (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

20 (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 10:

TGGAAAGTGT TGCCACAGGG CGCTGAAGCC TATCGCGTGC AGTTGCCGGA TGCCGCCTAT 60  
AGCCTCTACA TGGATGACAT CCTGCTGGCC TCC 93

25

(2) INFORMATION FOR SEQ ID NO: 11:

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 96 base pairs
- 30 (B) TYPE: nucleotide
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

35

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 11:

TTGGATCCAG TGYTGCCACA GGGCGCTGAA GCCTATCGCG TGCAGTTGCC GGATGCCGCC	60
TATAGCCTCT ACGTGGATGA CCTSCTGAAG CTTGAG	96

5

## (2) INFORMATION FOR SEQ ID NO: 12:

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 748 base pairs
- 10 (B) TYPE: nucleotide
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

## (ii) MOLECULE TYPE: cDNA

- 15 (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 12:

TGCAAGCTTC ACCGCTTGCT GGATGTAGGC CTCAGTACCG GN GTGCCCG CGCGCTGTAG	60
TT CGATGTAG AAAGCGCCCG GAAACACCGG GGACCAAATGC GT CGCCAGCT TG CGCGCCAG	120
20 CGCCTCGTTG CCATTGGCCA GCGCCACGCC GATATCACCC GCCATGGCGC CGGAGAGCGC	180
CAGCAGACCG GCGGCCAGCG GCGCATTCTC AACGCCGGC TCGTCGAACC ATT CGGGGGC	240
GATTCCGCA CGACCGCGAT GCTGGTTGGA GAGCCAGGCC CTGGCCAGCA ACTGGCACAG	300
GTTCAGGTAA CCCTGCTTGT CCCGCACCAA CAGCAGCAGG CGGGTCGGCT TGTCGCGCTC	360
GT CGTGATTG GTGATCCACA CGTCAGCCCC GACGATGGC TTCACGCCCT TGCCACGCC	420
25 TTCCCTGTAG ANGCGCACCA GCCCGAAGGC ATTGGCGAGA TCGGTCAGCG CCAAGGCGCC	480
CATGCCATCT TTGGCGGCAG CCTTGACGGC ATCGTCGAGA CGGACATTGC CATCGACGAC	540
GGAATATTGAG GAGTGGAGAC GGAGGTGGAC GAAGCGCGC GAATTCATCC GCGTATTGTA	600
ACGGGTGACA CCTTCCGCAA AGCATTCCGG ACGTGCCCGA TTGACCCGGA GCAACCCCGC	660
ACGGCTGCGC GGGCAGTTAT AATTCTGGCT TACGAATCAA CGGGTTACCC CAGGGCGCTG	720
30 AAGCCTATCG CGTGCAGTTG CCGGATGC	748

## (2) INFORMATION FOR SEQ ID NO: 13:

- 35 (i) SEQUENCE CHARACTERISTICS:  
(A) LENGTH: 18 base pairs

130

- (B) TYPE: nucleotide
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

5 (ii) MOLECULE TYPE: cDNA

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 13:

GCATCCGGCA ACTGCACCG

18

10

(2) INFORMATION FOR SEQ ID NO: 14:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 20 base pairs
  - (B) TYPE: nucleotide
  - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

20

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 14:

GTAGTTCGAT GTAGAAAGCG

20

25

(2) INFORMATION FOR SEQ ID NO: 15:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 18 base pairs
  - (B) TYPE: nucleotide
  - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

35

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 15:

131

GCATCCGGCA ACTGCACG

18

## 5 (2) INFORMATION FOR SEQ ID NO: 16:

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 23 base pairs
- (B) TYPE: nucleotide
- 10 (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

## 15 (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 16:

AGGAGTAAGG AAACCCAACG GAC

23

## 20 (2) INFORMATION FOR SEQ ID NO: 17:

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 19 base pairs
- (B) TYPE: nucleotide
- 25 (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

## 30 (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 17:

TAAGAGTTGC ACAAGTGCG

19

## 35 (2) INFORMATION FOR SEQ ID NO: 18:

132

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 21 base pairs
- (B) TYPE: nucleotide
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

5

## (ii) MOLECULE TYPE: cDNA

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 18:

10

TCAGGGATAG CCCCCATCTA T

21

## (2) INFORMATION FOR SEQ ID NO: 19:

15

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 24 base pairs
- (B) TYPE: nucleotide
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

20

## (ii) MOLECULE TYPE: cDNA

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 19:

25

AACCCCTTG CACTACATCA ATTT

24

## (2) INFORMATION FOR SEQ ID NO: 20:

30

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 15 base pairs
- (B) TYPE: nucleotide
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

35

133

(ii) MOLECULE TYPE: cDNA

(ix) FEATURES:

(B) LOCATION: 5, 7, 10, 13

5

(D) OTHER INFORMATION: G represents inosine (i)

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 20:

GGTCGTGCCG CAGGG

15

10

(2) INFORMATION FOR SEQ ID NO: 21:

(i) SEQUENCE CHARACTERISTICS:

15

(A) LENGTH: 21 base pairs

(B) TYPE: nucleotide

(C) STRANDEDNESS: single

(D) TOPOLOGY: linear

20

(ii) MOLECULE TYPE: cDNA

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 21:

TTAGGGATAG CCCTCATCTC T

21

25

(2) INFORMATION FOR SEQ ID NO: 22:

(i) SEQUENCE CHARACTERISTICS:

30

(A) LENGTH: 21 base pairs

(B) TYPE: nucleotide

(C) STRANDEDNESS: single

(D) TOPOLOGY: linear

35

(ii) MOLECULE TYPE: cDNA

134

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 22:

TCAGGGATAG CCCCCATCTA T

21

5

## (2) INFORMATION FOR SEQ ID NO: 23:

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 24 base pairs
- (B) TYPE: nucleotide
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

## (ii) MOLECULE TYPE: cDNA

15

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 23:

AACCCCTTG CACTACATCA ATTT

24

20

## (2) INFORMATION FOR SEQ ID NO: 24:

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 23 base pairs
- (B) TYPE: nucleotide
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

30

## (ii) MOLECULE TYPE: cDNA

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 24:

GCGTAAGGAC TCCTAGAGCT ATT

23

35

135

## (2) INFORMATION FOR SEQ ID NO: 25:

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 18 base pairs
- (B) TYPE: nucleotide
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

## (ii) MOLECULE TYPE: cDNA

10

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 25:

TCATCCATGT ACCGAAGG

18

15

## (2) INFORMATION FOR SEQ ID NO: 26:

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 20 base pairs
- (B) TYPE: nucleotide
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

## (ii) MOLECULE TYPE: cDNA

25

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 26:

ATGGGGTTCC CAAGTTCCCT

20

30

## (2) INFORMATION FOR SEQ ID NO: 27:

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 20 base pairs
- (B) TYPE: nucleotide
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

136

(ii) MOLECULE TYPE: cDNA

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 27:

5

GCCGATATCA CCCGCCATGG

20

(2) INFORMATION FOR SEQ ID NO: 28:

10

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 18 base pairs
- (B) TYPE: nucleotide
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

15

(ii) MOLECULE TYPE: cDNA

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 28:

20

GCATCCGGCA ACTGCACG

18

(2) INFORMATION FOR SEQ ID NO: 29:

25

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 20 base pairs
- (B) TYPE: nucleotide
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

30

(ii) MOLECULE TYPE: cDNA

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 29:

35

CGCGATGCTG GTTGGAGAGC

20

## (2) INFORMATION FOR SEQ ID NO: 30:

## 5 (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 20 base pairs
- (B) TYPE: nucleotide
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

10

## (ii) MOLECULE TYPE: cDNA

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 30:

15 TCTCCACTCC GAATATTCCG

20

## (2) INFORMATION FOR SEQ ID NO: 31:

## 20 (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 26 base pairs
- (B) TYPE: nucleotide
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

25

## (ii) MOLECULE TYPE: cDNA

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 31:

30 GATCTAGGCC ACTTCTCAGG TCCAGS

26

## (2) INFORMATION FOR SEQ ID NO: 32:

## 35 (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 23 base pairs

138

- (B) TYPE: nucleotide
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

5 (ii) MOLECULE TYPE: cDNA

## (ix) FEATURES:

- (B) LOCATION: 6, 12, 19
- (D) OTHER INFORMATION: G represents inosine (i)

10 (xi) SEQUENCE DESCRIPTION : SEQ ID NO: 32

CATCTGTTTG GGCAGGCAGT AGC

23

15 (2) INFORMATION FOR SEQ ID NO: 33:

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 24 base pairs
- (B) TYPE: nucleotide
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

25 (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 33:

CTTGAGCCAG TTCTCATACC TGGA

24

30 (2) INFORMATION FOR SEQ ID NO: 34:

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 22 base pairs
- (B) TYPE: nucleotide
- (C) STRANDEDNESS: single

139

(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

5 (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 34:

AGTGYTRCCM CARGGCCTG AA

22

10 (2) INFORMATION FOR SEQ ID NO: 35:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 22 base pairs
- (B) TYPE: nucleotide
- 15 (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

20 (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 35:

GMGGCCAGCA GSAKGTACATC CA

22

(2) INFORMATION FOR SEQ ID NO: 36:

25

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 22 base pairs
- (B) TYPE: nucleotide
- 30 (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 36:

35

GGATGCCGCC TATAGCCTCT AC

22

140

(2) INFORMATION FOR SEQ ID NO: 37:

**5 (i) SEQUENCE CHARACTERISTICS:**

- (A) LENGTH: 22 base pairs
  - (B) TYPE: nucleotide
  - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear

10

(ii) MOLECULE TYPE: cDNA

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 37:

15 AAGCCTATCG CGTGCAGTTG CC

22

(2) INFORMATION FOR SEQ ID NO: 38:

20

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 40 base pairs
  - (B) TYPE: nucleotide
  - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear

25

(ii) MOLECULE TYPE: cDNA

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 38:

30 TAAAGATCTA GAATTGGCT ATAGGGGGCA TCCGGGCAAGT

40

(2) INFORMATION FOR SEQ ID NO: 39

35

(i) SEQUENCE CHARACTERISTICS :

- (A) LENGTH : 50 amino acids

141

(B) TYPE : amino acid

(ii) MOLECULE TYPE : peptide

5 (xi) SEQUENCE DESCRIPTION : SEQ ID NO: 39

Asp Ala Phe Phe Cys Ile Pro Val Arg Pro Asp Ser Gln Phe Leu Phe  
1 5 10 15  
Ala Phe Glu Asp Pro Leu Asn Pro Thr Ser Gln Leu Thr Trp Thr Val  
10 20 25 30  
Leu Pro Gln Gly Phe Arg Asp Ser Pro His Leu Phe Gly Gln Ala Leu  
35 40 45  
Ala Gln  
50

15

(2) INFORMATION FOR SEQ ID NO: 40

(i) SEQUENCE CHARACTERISTICS :

- (A) LENGTH : 150 base pairs  
20 (B) TYPE : nucleic acid  
(C) STRANDEDNESS : single  
(D) TOPOLOGY : linear

(ii) MOLECULE TYPE : cDNA

25

(xi) SEQUENCE DESCRIPTION : SEQ ID NO: 40

GATGCCTTT TCTGCATCCC TGTACGTCCT GACTCTCAAT TCTTGTTCGC CTTTGAAGAT 60  
CCTTGAAACC CAACGTCTCA ACTCACCTGG ACTGTTTAC CCCAAGGGTT CAGGGATAGC 120  
30 CCCCCATCTAT TTGGCCAGGC ATTAGCCCAA 150

(2) INFORMATION FOR SEQ ID NO: 41

35 (i) SEQUENCE CHARACTERISTICS :

- (A) LENGTH : 11 amino acids

142

(B) TYPE : amino acid

(ii) MOLECULE TYPE : peptide

5 (xi) SEQUENCE DESCRIPTION : SEQ ID NO: 41

Cys Ile Pro Val Arg Pro Asp Ser Gln Phe Leu

1 5 10

10 (2) INFORMATION FOR SEQ ID NO: 42

(i) SEQUENCE CHARACTERISTICS :

(A) LENGTH : 17 amino acids

(B) TYPE : amino acid

15

(ii) MOLECULE TYPE : peptide

(xi) SEQUENCE DESCRIPTION : SEQ ID NO: 42

20 Val Leu Pro Gln Gly Phe Arg Asp Ser Pro His Leu Phe Gly Glu Ala

1 5 10 15

Leu

17

25

(2) INFORMATION FOR SEQ ID NO: 43

(i) SEQUENCE CHARACTERISTICS :

(A) LENGTH : 8 amino acid

30 (B) TYPE : amino acid

(ii) MOLECULE TYPE : peptide

(xi) SEQUENCE DESCRIPTION : SEQ ID NO: 43

35

Leu Phe Ala Phe Glu Asp Pro Leu

143

1                   5                   8

## (2) INFORMATION FOR SEQ ID NO: 44

5

## (i) SEQUENCE CHARACTERISTICS :

- (A) LENGTH : 8 amino acids
- (B) TYPE : amino acid

10                 (ii) MOLECULE TYPE : peptide

(xi) SEQUENCE DESCRIPTION : SEQ ID NO: 44

Phe Ala Phe Glu Asp Pro Leu Asn

15                 1                   5                   8

## (2) INFORMATION FOR SEQ ID NO: 45

20                 (i) SEQUENCE CHARACTERISTICS :

- (A) LENGTH : 25 base pairs
- (B) TYPE : nucleic acid
- (C) STRANDEDNESS : single
- (D) TOPOLOGY : linear

25

(ii) MOLECULE TYPE : cDNA

(xi) SEQUENCE DESCRIPTION : SEQ ID NO: 45

30   GTGCTGATTG GTGTATTTAC AATCC

25

## (2) INFORMATION FOR SEQ ID NO: 46

35                 (i) SEQUENCE CHARACTERISTICS :

- (A) LENGTH : 1859 base pairs

- (B) TYPE : nucleic acid
- (C) STRANDEDNESS : single
- (D) TOPOLOGY : linear

5 (ii) MOLECULE TYPE : cDNA

(xi) SEQUENCE DESCRIPTION : SEQ ID NO: 46

	G TGCTGATTG GTGTATTTAC AATCCTTTAT CTAATCCGAA ATGCCCATGT TGCAATATGG	60
10	AAAGAAAGGG AGTTCCTAAC CTCTGGGGGA ACCCCCATTAA ATACCAACAA GTAAATCATG	120
	GAGTTATTGC ACACAGTGCA AAAACTCAAG GAGGTGGAAG TCTTACACTG CCAAAGCCAT	180
	CAGAAAAGGG AAGAGGGGAG AAGAGCACCA TAAGTGGCTA CAGAGGCAAG GAAAGACTAG	240
	CAGAAAGGAA AGAGAGAAAG AGACAGAAAG TCAGAGAGAG AGAGAGGAAG AGACAGAGCA	300
	CAAAGAGGGA GTCAGAGAGA GAGAGAGACA GAGAGTCAGA GAGAAGGAAA GAGAGAGAGG	360
15	AAGAGACAAA GAATGAATCA AACAGAGAGA CAGAAAGTCA GAGAGAGAGA GAGAGAGGAA	420
	GAGACAGAGA AAAAGAGGGG GTCAGAAAAA GAGAGACCAA AGAAGAAGTC CAAAGAGAAA	480
	GAAAGAGAGA TGGAAGTAGT AAAGGAAAAA CAGTGTACCC TATTCTTTA AAAGCCGGGG	540
	TAAATTAAA ACCTATAATT GATAACTGAA GGTCTTCTCT GTAACCCTGT AACACTCCAA	600
	TACCACTTG TTGTCAAGTG TAAACAAGGG CGTAGCCAA AAGCACTGAG GCCACTAACAA	660
20	ACCCATAGCC TTCCTATCAA AATTCTTAA CCCAGCAGGT TTCTAACAG GGGATCTAAA	720
	TCTTAATTAA TTACCATACA ATGGTCCAAC CAGACTTAGG AGGAATTCCC TTCAGGACGG	780
	GAAGATAGAT GCTTCCTCCC AGCGATTAA GGGAGAAAGA CACAATGGGT ATTCAAGTAAG	840
	TGCCAAGGGG AACACTTGTA GAAGCAAAGT TAGGAAAATT GCCAAATAAT TGGTTGCTC	900
	AAGAGTTGTT TGCACTCAGC CAAACCTTGA AGTACTTGCA GAATCAGAAA GGAGCCATCT	960
25	ATACCAATTC TAAGTTAATA TGGACTGAAG GAGGTTTAT TAATACCAAA GAGAAATTAA	1020
	AATCCCAAAC TTATAAGGTT TTCAACCCAA GTAAAGTTG CTAAAAGTTA ACAGCGTAAC	1080
	ATGTATTATC CTACTACCAC ACACTCTCAA AGGATTCTC AGACAGTTG CAAGAAATAA	1140
	TGATATCTAT CCTTACTCTA CAATCCAAA TAGACTCTT GGCAGCAGTG ACTCTCCAAA	1200
	ACCGTCAAGG CCTAGACCTC CTCACTGCTG AGAAAGGAGG ACTCTGCACC TTCTTAAGGG	1260
30	AAGAGTGTG TCTTTACACT AACCAAGTCAG GGATAGTATG AGATGCTGCC CGGCATTTAC	1320
	AGAAAAAGGC TTCTGAAATC AGACAACGCC TTTCAAATTC CTATACCAAC CTCTGGAGTT	1380
	GGGCAACATG GTTTCTTCCC TTTCTATGTC CCATGGCTGC CATCTTGCTA TTACTCGCCT	1440
	TTGGGCCCTG TATTTTAAAC CTCCCTGTCA AATTGTTTC TTCTAGGATC GAGGCCATCA	1500
	AGCTACAGAT GGTCTTACAA ATGGAACCCC AAATGAGCTC AACTATCAAC TTCTACTGAG	1560
35	GACCCCTAGA CCAACCCCCCT GGCCCTTCA CTGGCCTAAA GAGTTCCCT CTGGAGGACA	1620
	CTACCACTGC AGGGCCCCAT CTTGCCCCCT ATCCAGAAGG AAGTAGCTAG AGCAGTCATT	1680

145

GCCCAATTCC CAAGAGCAGC TGGGGTGTCC CGTTTAGAGT GGGGATTGAG AGGTGAAGCC 1740  
AGCTGGACTT CTGGGTGGG TGGGGACTTG GAGAACTTT GTGTCTAGCT AAAGGATTGT 1800  
AAATGCAACA ATCAGTGCTC TGTGTCTAGC TAAAGGATTG TAAATACACC AATCAGCAC 1859

5

## (2) INFORMATION FOR SEQ ID NO: 47

## (i) SEQUENCE CHARACTERISTICS :

- 10 (A) LENGTH : 23 base pairs  
(B) TYPE : nucleic acid  
(C) STRANDEDNESS : single  
(D) TOPOLOGY : linear

15 (ii) MOLECULE TYPE : cDNA

(xi) SEQUENCE DESCRIPTION : SEQ ID NO: 47

TGATGTGAAC GGCATACTCA CTG

23

20

## (2) INFORMATION FOR SEQ ID NO: 48

## (i) SEQUENCE CHARACTERISTICS :

- 25 (A) LENGTH : 24 base pairs  
(B) TYPE : nucleic acid  
(C) STRANDEDNESS : single  
(D) TOPOLOGY : linear

30 (ii) MOLECULE TYPE : cDNA

(xi) SEQUENCE DESCRIPTION : SEQ ID NO: 48

CCCAGAGGTT AGGAACCTCCC TTTC

24

35

146

## (2) INFORMATION FOR SEQ ID NO: 49

## (i) SEQUENCE CHARACTERISTICS :

- (A) LENGTH : 25 base pairs
- (B) TYPE : nucleic acid
- (C) STRANDEDNESS : single
- (D) TOPOLOGY : linear

## (ii) MOLECULE TYPE : cDNA

10

## (xi) SEQUENCE DESCRIPTION : SEQ ID NO: 49

GCTAAAGGAG ACTTGTGGTT GTCAG

25

15

## (2) INFORMATION FOR SEQ ID NO: 50:

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 22 base pairs
- (B) TYPE: nucleotide
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

## (ii) MOLECULE TYPE: cDNA

25

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 50:

CAACATGGGC ATTTCGGATT AG

22

30

## (2) INFORMATION FOR SEQ ID NO: 51:

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 400 base pairs
- (B) TYPE: nucleotide
- (C) STRANDEDNESS: single

35

## (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

5 (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 51:

GGCTGCTAAA	GGAGACTTGT	GGTTGTCAGA	CAATGCCCTA	CTTAGGTACC	AGGCCTTATT	60
ACTTGAGGGA	CTGGTGCTTC	AGATGCGCAC	TTGTGCAGCT	CTTAACCCAA	ACTTATGCTG	120
CCCAGAACGA	TCTTTAGAG	GTCCCCCTAG	CCAACCTGA	CCTCAACCTA	TATATATACT	180
10 GATGGAAGTT	CGTTGTAGA	AAAGGGATTA	CAAAGGGNAG	GATATNCCAT	AGGTTAGTGA	240
TAAAGCAGTA	CTTGAAAGTA	AGCCTCTTCC	CCCCAGGGAC	CAGCGCCCCC	GTTAGCAGAA	300
CTAGTGGCAC	TGACCCCGAG	CCTTAGAACT	TGGAAAGGGA	GGAGGATAAA	TGTGTATACA	360
GATAGCAAGT	ATGCTTATCT	AATCCGAAAT	GCCCCATGTTG			400

15

(2) INFORMATION FOR SEQ ID NO: 52:

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 2389 base pairs
- 20 (B) TYPE: nucleotide
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

25

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 52:

TCAGGGATAG	CCCCCATCTA	TTGGTCAGG	CACTGGCCCA	AGATCTAGGG	ACATGCCACT	60
TTTAAGAGCC	ATTTCTCAAG	TCCAGGTACT	CTGGTCCTTC	GGTATGTGGA	TGATTTACTT	120
30 TTGGCTACCA	GTTCACTAGC	CTCATGCCAG	CAGGCTACTC	TAGATCTCTT	GAACCTTCTA	180
GCTAATCAAG	GGTACAAGGC	ATCTAGGTTG	AAGGCCAGC	TTGCCTACA	GCAGGTCAA	240
TATCTAGGCC	TAATCTTAGC	CAGAGGGACC	AGGGCACTCA	GCAAGGAACA	AATACAGCCT	300
ATACTGGCTT	ATCCTCACCC	TAAGACATTA	AAACAGTTGC	GGGGGTTCCCT	TGGAATCACT	360
GGCTTTTGG	TGACTATGGA	TTCCCGAGATA	CAGCAAGATT	GGCAGGCC	TCTATACTGT	420
35 AATCAAGGAG	ACTCACGAGG	GCAAGTACTC	ATCTAGTACA	ATGGGAACTA	GGGACAGAAA	480
CAGCCTTCAA	AACCTTAAAG	CAGGCCCTAG	TACAATCTCC	ACCTTTAACG	CTTCCCACAG	540

GACAAAAC	TTT	CTT	TAT	AAC	AGA	GAG	AGA	GAG	ATC	ACAGA	GAG	GGG	CAG	AGA	GT	C	CTT	GGG	T	TTC	ATT	C			
AGACTCAT	GG	GA	CT	AC	CC	CA	CC	AG	TG	GA	CA	CC	TA	AG	AA	TT	AT	GT	GT	AG	TA	GT	AG		
CAA	AA	GG	CT	GC	CT	CA	CT	GT	GG	GT	GT	CT	TT	GT	CA	AG	AA	GG	CT	GT	CA	AG	AA		
CTATCA	AA	AA	AT	AA	AA	AT	AA	AA	GG	AT	CT	CA	GG	AC	T	AC	T	CA	TG	AT	GA	AT	GG		
5	ACTAGGTGCC	AAA	AGA	AG	TT	TAT	GGG	TAT	TC	AGA	CA	ACC	AC	CT	GCT	TTA	GA	GGG	ACT	GGG	ACT	GGG	ACT	GGG	
ACTC	CT	GG	GAG	GAT	GCT	CA	AG	TG	CG	TT	TT	GT	GG	CC	TCA	AC	CC	CT	GC	CA	CT	TT	CC	CT	
CC	AG	AG	GAT	G	G	G	AG	AG	CC	GC	TG	AC	AT	GC	TG	CC	AA	TT	AT	TC	TT	CC	TT	CC	
AC	CC	GAG	AT	G	AT	T	CT	TT	AG	TA	CC	CT	TA	AT	CC	TA	CT	AA	AC	CT	TA	AC	CA	AT	
GA	AG	TT	CATT	TG	TG	AA	AA	AC	GG	AT	AT	GAA	GG	CAG	GT	TA	AG	TT	AG	TG	AT	GT	AA	1080	
10	TCATA	ACT	TG	C	A	AG	T	AA	GG	C	CT	CC	AG	GC	AC	TC	AG	TT	AG	CA	GA	AC	TA	GT	CA
CA	CT	TAC	CT	TT	AA	CT	TT	AA	GG	GG	GG	AA	AG	AG	AC	TC	AA	CT	AC	CT	TA	AC	CA	AT	AT
AG	TAT	G	CT	TA	AA	T	CT	AC	CC	AT	CC	AT	AT	GC	T	GA	GG	AA	AG	G	AT	TC	CT	AA	1140
CC	CC	TGG	GG	GG	GG	GG	GG	GG	GG	GG	GG	GG	GG	GG	GG	GG	GG	GG	GG	GG	GG	GG	GG	GG	GG
CA	AA	AA	CT	CA	AA	AA	CT	CA	AA	AA	AA	AA	AA	AA	AA	AA	AA	AA	AA	AA	AA	AA	AA	AA	AA
15	GGG	AGA	AC	AG	C	AG	C	AT	AG	GT	GG	CAG	CAG	AC	TC	AG	TT	AA	GG	GA	AG	GG	GA	AG	AG
GAG	ACA	AC	GT	CA	AC	GAC	AG	AA	AG	GG	AA	AG	AG	AG	AC	AG	AA	AG	AG	AG	AG	AG	AG	AG	1380
AC	AG	TT	AG	TC	CA	AG	AG	AG	AG	AG	AG	AG	AG	AG	AC	AG	AA	AG	AG	AG	AG	AG	AG	AG	1500
GG	AA	AG	AG	AG	GA	AG	AG	AC	CC	NA	AG	AG	AA	AA	GAG	AT	AG	AA	AG	AG	AA	AG	AG	AG	1620
AA	AA	AC	AT	TG	TT	AA	TT	AA	AG	CC	TT	AA	AC	CT	TA	AA	AC	CT	TA	AA	AC	CT	TA	AA	1680
20	TGAG	TT	CT	TG	CA	CC	CT	CT	CC	GG	GG	GG	AT	Y	CT	GG	GG	GG	GG						
AA	TT	GT	GG	GT	CG	TC	CC	CT	AT	TC	TA	CC	AT	AC	CC	TT	GT	TT	AG	GT	GG	AA	TT	GT	AA
CG	AG	GG	GT	TA	GAG	CC	AG	GG	AC	CT	GAC	AA	TC	CC	TT	TC	TA	CC	AA	AT	TC	CC	AA	AT	1860
CCT	TA	AC	CC	CA	GC	AG	TT	TT	TC	TAAA	AG	GG	GA	T	CT	AA	AT	CT	TT	AA	AC	CC	AA	AG	1920
TCA	AA	AC	CAG	AG	TCT	AG	GG	AG	GA	GG	AC	GG	AT	GT	GG	TT	CT	CC	AG	GC	AG	GC	AG	GC	1980
25	GAT	AA	AG	AA	AA	AA	AA	AA	AA	AA	AA	AA	AA	AA	AA	AA	AA	AA	AA						
AC	CA	GT	TT	AG	GG	CC	TA	AA	TT	GG	TCT	ACT	CC	AA	AT	GT	GT	GAG	T	TG	TG	CA	CT	AC	2040
TA	AT	AT	GG	AC	AG	AG	AG	AG	AG	AG	AG	AG	AG	AG	AC	AG	AA	AG	TC	CC	CA	CT	AC	2100	
TA	AT	AT	GG	AC	AG	AG	AG	AG	AG	AG	AG	AG	AG	AG	AC	AG	AA	AG	AG	AG	AG	AG	AG	AG	2160
GG	TT	TT	CA	AC	TAA	AG	AA	AT	TT	TA	TA	TA	TA	TA	TA	TA	TA	AA	AC	CT	CT	AC	CA	2220	
GG	TT	TT	CA	AC	TAA	AG	AA	AT	TT	AC	TA	AA	AG	AA	AG	AT	AC	CT	CT	AC	CA	2280			
30	CA	AC	AC	AC	TC	AN	AG	GG	AT	C	CT	CAG	AC	AG	TT	T	T	T	AC	AA	AG	AA	AT	AC	AA
AG	GA	TAG	TAA	CT	AC	AA	AT	CC	CC	AA	AT	AC	AT	TC	TC	TA	TT	GG	CAG	CA	GC	TG	AC	TCT	2340
																								2389	

(2) INFORMATION FOR SEQ ID NO: 53:

35

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 2448 base pairs
- (B) TYPE: nucleotide
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

5

(ii) MOLECULE TYPE: cDNA

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 53:

10	TCAGGGATAG CCCCCATCTA TTTGATCAGG CACTAGCCCA AGATCTAGGC CACTTCTGAA	60
	GTCCAGGCAT TCTAGTCCTT CAGTATGTGG ATGATTTACT TTTGGCTACC AGTTTCCAAG	120
	CCTCATGCCA GCAGGCTACT TGAGATCTCT TGAACCTTCT AGCTAATCAA GGGTGTATGG	180
	CATCTAAATT GAAAGTCCAG CTCTGCCTAC AACAAAGTCAA ATATCTAGGC CTAATCTTAG	240
	ATAGAAGAAC CAGGGCCCTC AGCAAGGAAT GAATAAAGCC TATGCTGGCT TATCGGCACC	300
15	CTAAGACATT AAAACAATTG TGGGGGTTCC TTGGAATCAC TGGCTTTGC CGACTATGGA	360
	TCCCTGGATA GAGTGAGATA GCCAGGCCCT CTCTATTACT CTTATCAAGG AGACCCAGAG	420
	GGCAAATACT TATCTAGTAT TATGGGNACC AGAGGCAGAA AAAGCCTTCC AAACCTTAAA	480
	GGAGACCCCTA GTACAAGCTC CAGCTTAAG CCTTCCCACA GGACAAANCT TCTCTTTATA	540
	TGTCACAGAG AGAGCAGGAA TAGCTCCTGG AGTCCTTACT CAGACTTTG GACGACCCCA	600
20	CGGCCAGTGG CRTACCTAAG TAAGGAAATT GATGTAGTAG CAAAAGGCTG GCCTCACTGT	660
	TTATGGGTAG TTGCGGCTGT GGCAGTCTTA CTGTCAAAGG CTATCAAAT AATACAAGGA	720
	AAGGATTTCAGT CTATCTGGAC TACTCATGAG GAAAATGGCA TATTAGGTGC CAAAGGAAGT	780
	TTTTGGCTAT CAGACAACCA CCTGCTCAGA TTCCAGGCAC TACTGATTGA GAGACCAGTG	840
	CTTTAAATAT GTATGTGTGT GTGTGGCCCT CAACCCTGCC ACTGTTCTCC CAGAAGATGG	900
25	AGAACCAATG AAGCATTACT GTCAACAAAT TAGAGTCCAG AGTTATGCTG CCTGAGAGGA	960
	TCTCTTAGAA GTCCCCCTAG CTAATCCTGA CCTTAACCTA TATGCTGATG GAAGTCACT	1020
	TGTGGAGAAT GGGATACGAA AAGCACATTA TGCCATAGTT AGTGAGGTAA CAGTACTTGA	1080
	AAGTAAGCCT ATTCCCCCAT GGACCAGAGC CCAGTTAGCA GAACTAGTGG CACTTACCCA	1140
	AGCCTTAGAA CTAGGAAAGG GAAAAATAAT AAATGTGTAT ACAGATAGCA AGTATGCTTA	1200
30	TCTAATCCTA CATGCCCATG CTGCAGTATG GAAAGAAAGG GAGTTCTAA CCTCTGGGG	1260
	AACCCCCATT AAATACCACA AGGCAAATCA TGGAGTTATT GCATGTAGTG CAAAACCTCA	1320
	AGTAGGTGGC AGTTTACAC TGCCTGAAGC TATGGGAAG GAGAGAGGAG AACAGCAGCA	1380
	TAAGTGGCTA GCAGAGGCAG CGAAAGACTA GCAGAGAGGA GAGGTAGGGG AAAGACAGAA	1440
	AGTCAAAGAA AAGAAGTCAA AGACAGACAG AGAAAGAGAC AGAGGGAGCC AGAGAGAAAG	1500
35	AAAAGAGAGA ACGAAAGAGA CAGAATGTCA AAGAACAGAA GAGAGAGGCA GCGCCAGAAG	1560
	AGTTAAGAAA GTGAGAAAGA GAGATGGAAA TAGTAAAGAA AAAACAGTGT ACCCTATTCC	1620

150

TTTAAAAGCC AGGGTAAATT TAAAACGTAT AATTTATAA TTGGAAGGTC TTCTCCATAA 1680  
CCCTATAACA TTAAAATACC ACCTTGTGTC CAGTGTAAAC AAGAGCATAG CCCAAAAGCA 1740  
CTGAGGCCAC TGACAACCCA TAGCCTTCCT ATCAAAAATC CTTAACTCTG CAGGTTTCCT 1800  
AACAGGGGAT CTAAATCTCA ACTAATCACC ATACAATGGT CCGACCAGAC CTAGGAGCGA 1860  
5 CTCCCCCTCAG GACAGAAGGA TGGATGGTTC CTCCCAGGCC ATTAAGGGAA AGAGACACAA 1920  
TGGGTATTCA GTAACTGATA AGGGAACACTCT TGTAGAAGCA GTTAGGAAGA TTGCCTAATA 1980  
TTTGGTCTGC TCAAATGTGC CAGCTGTTG CACTCAGCTA AACCTTAAAT TACTTACAGA 2040  
ATTAGGAAGG AGCCATCTAT ACCAATTCTG AGTTAATATG AGCTGAACAA GTTCTTATTA 2100  
ATACCAAAGA ATCATTGAAA TCTCAAACCTT GCAGGTTTT CAACAAAAGT AAAGTTTGCT 2160  
10 GAAAGTTAGC AGTGTAAACAT GTATTATCCT AACTTCTAAT CTTGTGGAAA TCAGACCCCTA 2220  
TCAGTGCCCC TCAAAGCTGA AGTCCATCAG CATATGGCCA TACAACCTAAT ACCCCTATTT 2280  
ATAGGGTTAG GAATGGCCAC TGCTACAGGA ATGGGAGTAA CAGGTTTATC TACTTCATTA 2340  
TCCTATTACC ACACACTCTT AAAGGATTTC TCAGACAGTT TACAAGAAAT AACAAAATCT 2400  
ATCCTTACTC TNTARTCCCA AATAGRTTCT TTGGCAGCAG TGACTCTC 2448

15

## (2) INFORMATION FOR SEQ ID NO: 54:

## (i) SEQUENCE CHARACTERISTICS:

- 20 (A) LENGTH: 21 base pairs  
(B) TYPE: nucleotide  
(C) STRANDEDNESS: single  
(D) TOPOLOGY: linear

25 (ii) MOLECULE TYPE: cDNA

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 54:

CCTGAGTTCT TGCACCTAACCC C

21

30

## (2) INFORMATION FOR SEQ ID NO: 55:

## (i) SEQUENCE CHARACTERISTICS:

- 35 (A) LENGTH: 23 base pairs  
(B) TYPE: nucleotide

151

(C) STRANDEDNESS: single

(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

5

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 55:

GTCCGTTGGG TTTCCTTACT CCT

23

10

(2) INFORMATION FOR SEQ ID NO: 56:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 1196 base pairs

15

(B) TYPE: nucleotide

(C) STRANDEDNESS: single

(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

20

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 56:

	TTCCTGAGTT CTTGCACTAA CCTCAAATGA GAGAAGTGCC GCCATAACTG CAACCCAAGA	60
	GTTTGGCGAT CCCTGGTATC TCAGTCAGGT CAATGACAGG ATGACAACAG AGGAAAGATA	120
25	ATGATTCCCC ACAGGCCAGC AGGCAGTTCC CAGTGTAGAC CCTCATTAGG ACACAGAAC	180
	AGAACATGGA GATTGGTGCC GCAGACATTT GCTAACTTGC GTGCTAGAAC GACTAAGGAA	240
	AACTAGGAAG ATATGAATTA TTCAATGATG TCCACTATAA CACAGGGAA AGGAAGAAAA	300
	TCCTACTGCC TTTCTGGAGA GACTAAGGGA GGCATTGAGG AAGCATAACCA GGCAAGTGG	360
	CATTGGAGGC TCTGGAAAAG GGAAAAGTTG GGAAAAGTAT ATGTCTAATA GGGCTTGCTT	420
30	CCAGTGTGGT CTACAAGGAC ACTTTAAAAA AGATTGTCCA ATAGAAATAA GCCACCACCT	480
	CGTCCATGCC CCTTATGTCA AGGGAATCAC TGGAAGGCC ACTGCCAGG GGGATGAAGG	540
	TCCTCTGAGT CAGAAGCCAC TAACCAGATG ATCCAGCAGC AGGACTGAGG GTGCCGGGG	600
	CAAGCGCCAG CCCATGCCAT CACCCCTACA GAGCCCCAGG TATGCTTGAC CATTGAGGGT	660
	CAGAAGGGTA CTGTCTCCTG GACACTGGCG GGCCTTCTCA GTCTTACTTT CCTGTCCCTGG	720
35	ACAACTGTCC TCCAGATCTG TCACTGTCCG AGGGGTCTA GGACAGCCAG TCACTAGATA	780
	CTTCTCCCAG CCACTAAGTT GTGACTGGGG AACTTACTC TTCCACATGC TTTCTAATT	840

ATGCCTGAAA GCCCCACTCT CTTGTTAGGG GAGAGACATT CTAGCAAAAG CAGGGGCCAT 900  
TATACATGTG AATATAGGAG AAGGAACAAC TGTTGTTGT CCCCTGCTTG AGGAAGGAAT 960  
TAATCCTGAA GTCCGGGCAA CAGAAGGACA ATATGGACAA GCAAAGAATG CCCGTCCTGT 1020  
TCAAGTTAAA CTAAAGGATT CCACCTCCTT TCCCTACCAA AGGCAGTACC CCCTCAGACC 1080  
5 CGAGACCCAA CAAGAACTCC AAAAGATTGT AAAGGACCTA AAAGCCCAAG GCCTAGTAAA 1140  
ACCAAGCAAT AGCCCTTGCA AGACTCCAAT TTTAGGAGTA AGGAAACCCA ACGGAC 1196

## (2) INFORMATION FOR SEQ ID NO: 57:

10

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 2391 base pairs
- (B) TYPE: nucleotide
- (C) STRANDEDNESS: single
- 15 (D) TOPOLOGY: linear

## (ii) MOLECULE TYPE: cDNA

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 57:

20

ATGATCCAGC AGCAGGACNG AGGGTCCCCG GGGCAAGCGC CAGCCCATGC CATCACCCCTC 60  
ACAGAGCCCC AGGTATGCTT GACCATTGAG GGTCAGAAGG GTNACTGTCT CCTGGACACT 120  
GGCGGNGCCT TCTCAGTCTT ACTTTCTGT CCTGGACAAC TGTCCCTCAG ATCTGTCACT 180  
GTCCGAGGGG TCCTAGGACA GCCAGTCACT AGATACTTCT CCCAGCCACT AAGTTGTGAC 240  
25 TGGGGAACTT TACTCTTCCC ACATGCTTT CTAATTATGC CTGAAAGCCC CACTCTCTG 300  
TTGGGGAGAG ACATTCTAGC AAAAGCAGGG GCCATTATAC ATGTGAATAT AGGAGAAGGA 360  
ACAACTGTTT GTTGTCCCC GCTTGAGGAA GGAATTAATC CTGAAGTCCC GGCAACAGAA 420  
GGACAATATG GACAAGCAAA GAATGCCGT CCTGTTCAAG TTAAACTAAA GGATTCACC 480  
TCCTTCCCT ACCAAAGGCA GTACCCCTC AGACCCGAGA CCCAACAAAGA ACTCCAAAAG 540  
30 ATTGTAAAGG ACCTAAAAGC CCAAGGCCTA GTAAAACCAA GCAATAGCCC TTGCAAGACT 600  
CCAATTTAG GAGTAAGGAA ACCCAACGGA CAGTGGAGGT TAGTGCAAGA ACTCAGGATT 660  
ATCAATGAGG CTGTTGTTCC TCTATACCA GCTGTACCTA ACCCTTATAC AGTGCTTCC 720  
CAAATACCAAG AGGAAGCAGA GTGGTTACA GTCCTGGACC TTAAGGATGC CTTTTCTGC 780  
ATCCCTGTAC GTCCTGACTC TCAATTCTG TTTGCCTTG AAGATCCTTT GAACCCAACG 840  
35 TCTCAACTCA CCTGGACTGT TTTACCCAA GGGTCAGGG ATAGCCCCCA TCTATTGGC 900  
CAGGCATTAG CCCAAGACTT GAGTCAATT TCATACCTGG ACACTCTTGT CCTTCAGTAC 960

ATGGATGATT TACTTTAGT CGCCC GTTCA GAAAC CTTGT GCCAT CAAGC CACCC AAGAA 1020  
CTCTTAAC TT CACTAC CTG TGCTAC AAG GTTCCA AACCAA AGGC TCGGCTCTGC 1080  
TCACAGGAGA TTAGATACTN AGGGCTAAAA TTATCCAAAG GCACCAGGGC CCTCAGTGAG 1140  
GAACGTATCC AGCCTATACT GGCTTATCCT CATCCCCAAA CCTCTAAAGCA ACTAAGAGGG 1200  
5 TTCCTGGCA TAACAGGTTT CTGCCGAAAA CAGATTCCCA GGTACASCCC AATAGCCAGA 1260  
CCATTATATA CACTAATTAN GGAAACTCAG AAAGCCAATA CCTATTTAGT AAGATGGACA 1320  
CCTACAGAAG TGGCTTTCCA GGCCCTAAAG AAGGCCCTAA CCCAAGCCCC AGTGTTCAGC 1380  
TTGCCAACAG GGCAAGATTT TTCTTTATAT GCCACAGAAA AAACAGGAAT AGCTCTAGGA 1440  
GTCCTTACGC AGGTCTCAGG GATGAGCTTG CAACCCGTGG TATACTGAG TAAGGAAATT 1500  
10 GATGTAGTGG CAAAGGGTTG GCCTCATNGT TTATGGTAA TGGNGGCAGT AGCAGTCTNA 1560  
GTATCTGAAG CAGTTAAAAT AATACAGGGA AGAGATCTTN CTGTGTGGAC ATCTCATGAT 1620  
GTGAACGGCA TACTCACTGC TAAAGGAGAC TTGTGGTTGT CAGACAACCA TTTACTTAAN 1680  
TATCAGGCTC TATTACTTGA AGAGCCAGTG CTGNGACTGC GCACCTGTGC AACTCTTAAA 1740  
CCCAAACCTTA TGCTGCCAG AAGGATCTTT NTAGAGGTCC CCTTAGCCAA CCCTGACCTC 1800  
15 AACTATATAT ATACTGATGG AAGTCGTTT GTAGAAAAGG GATTACAAAG GGNAGGATAT 1860  
NCCATAGGTG TTAGTGATAA AGCAGTACTT GAAAGTAAGC CTCTTCCCCC CCAGGGACCA 1920  
GCGCCCCCGT TAGCAGAACT AGTGGCACTG ACCCCCGCAG CCTTAGAACT TTGGAAAGGG 1980  
AGGAGGATAA ATGTGTATAC AGATAGCAAG TATGCTTATC TAATCCGAAA TGCCCATGTT 2040  
GTTTATCTAA TCCGAAATGC CCATGTTGCA ATATGGAAAG AAAGGGAGTT CCTAACCTCT 2100  
20 GGGGAACCC CCATTAATAA CCACAAGTTA ATCATGGAGT TATTGCACAC AGTGCAAAAA 2160  
CTCAAGGAGG TGGAAAGTCTT ACAC TGCCAA AGCCATCAGA AAAGGGAAAG GGGAGAAGAG 2220  
CAGCATAAGT GGCTACAGAG GCAAGGAAAG ACTAGCAGAA AGGAAAGAGA GAAAGAGACA 2280  
GAAAGTCAGA GAGAGAGAGA GGAAGAGACA GAGCACAAAG AGGGAGTCAG AGAGAGAGAG 2340  
AGACAGAGAG TCAGAGAGAA GGAAAGAGAG AGAGGAAGAG ACAAAAGAATG A 2391

25

## (2) INFORMATION FOR SEQ ID NO: 58:

## (i) SEQUENCE CHARACTERISTICS:

- 30 (A) LENGTH: 1722 base pairs  
(B) TYPE: nucleotide  
(C) STRANDEDNESS: single  
(D) TOPOLOGY: linear

35 (ii) MOLECULE TYPE: cDNA

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 58:

	TGGAGAATAG CAGCATAAGT TGGCTGGCAG AAGTAGGGAA AGACAGCAAG AAGTAAAGAA	60
	AAAAARGAGA AAGTCAGAGA AAGAAAAAAA GAGAGGAAGA AACAAAGAAG AACTTGAAGA	120
5	GAGAAAGAAG TAGTAAAGAA AAAACAGTAT ACCCTATTCC TTTAAAAGCC AGGCTAAATT	180
	TCTGTCTACC TAGCCAAGGC ATATTCTTCT TATGTGGAAC ATCAACCTAT ATCTGCCCTCC	240
	CCACTAACTG GACAGGCACC TGAACCTTAG TCTTTCTAAG TCCTAACATT AACATTGCC	300
	CAGGAAATCA GACCCTATTG GTACCTGTCA AAGCTAAAGT CCCGTCAGTG CAGAGCCATA	360
	CAACTAATAT CCCTATTAT AGGGTTAGGA ATGGCTACTG CTACAGGAAC TGGAAATAGCC	420
10	GGTTTATCTA CTTCATTATC CTACTACCAT ACACCTCTAA AGAATTCTC AGACAGTTG	480
	CAAGAAATAA TGAAATCTAT TCTTACTTTA CAATCCCAAT TAGACTCTT GGCAGCAATG	540
	ACTCTCCAAA ACCGCCGAGG CCCACACCTC CTCACTGCTG AGAAAGGAGG ACTCTGCACC	600
	TTCTTAGGGG AAGAGTGTG TTTTACACT AACCAAGTCAG GGATAGTACG AGATGCCACC	660
	TGGCATTAC AGGAAAGGGC TTCTGATATC AGACAATGCC TTCAAACCTC TTATACCAAC	720
15	CTCTGGAGTT GGGCAACATG GCTTCTTCCA TTTCTAGGTC CCATGGCAGC CATCTTGCTG	780
	TTACTCACCT TTGGGCCCTG TATTTTAAG CTTCTTGTCA AATTGTTTC CTCTAGGATC	840
	GAAGCCATCA AGCTACAGAT GGTCTTACAA ATGGAACCCC AAATGAGTTC AACTAACAAAC	900
	TTCTACCAAG GACCCCTGGA ACGATCCACT GGCACTTCCA CTAGCCTAGA GATTCCCCCTC	960
	TGGAAGACAC TACAACGTCA GGGCCCTTC TTGCCCCCTA TCCAGCAGGA AGTAGCTAGA	1020
20	GCGGTCATCG GCCAAATTCC CAACAGCAGT TGGGGTGTCC TGTTAGAGG GGGGATTGAA	1080
	GAGGTGACAG CCTGCTGGCA GCCTCACAGC CCTCGTTGGY TCTCAGTGCC TCCTCAGCCT	1140
	TGGTGCCAC TCTGGCCGTG CTTGAGGAGC CCTTCAGCCT GCCACTGCAC TGTGGGAGCC	1200
	TCTTTCTGGG CTGGACAAGG CCGGAGCCAG CTCCCTCAGC TTGCAGGGAG GTATGGAGGG	1260
	AGAGATGCAG GCGGGAACCA GGGCTGCGCA TGGCGCTTGC GGGCAGCAT GAGTTCCAGG	1320
25	TGGGCGTGGG CTCGGCGGGC CCCACACTCG GGCAGTGAGG GGCTTAGCAC CTGGGCCAGA	1380
	CAGATGCTGT GCTCAACTTC TTGCTGGGC CTTAGCTGCC TTCCCCGTGG GGCAGGGCTY	1440
	CGGGAACMTG CAGCCTGCC ATGCTTGAGC CCCCCACCCC GCCGTGGTT CYTGCACAGC	1500
	CCAAGCTTCC CGGACAAGCA CCACCCCTTA TCCACGGTGC CCAGTCCCCT CAACCACCCA	1560
	AGGGTTGAGG AGTGCAGGCA CACAGCGCGG GATTGGCAGG CAGTTCCACT TGCAGGCTTG	1620
30	GTGCGGGATC CACTGCGTGA AGCCAGCTGG GCTCCTGAGT CTGGTGGGGA CTTGGAGAAT	1680
	CTTTATGTCT AGCTAAGGGA TTGTAAATAC ACCAATCAGC AC	1722

## (2) INFORMATION FOR SEQ ID NO: 59:

155

- (A) LENGTH: 495 base pairs
- (B) TYPE: nucleotide
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

5

(ii) MOLECULE TYPE: cDNA

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 59:

10 CTTCCCCAAC TAATAAGGAC CCCCTTTCA ACCCAAACAG TCCAAAAGGA CATAGACAAA 60  
GGAGTAAACA ATGAACCAAA GAGTGCCAAT ATTCCCTGGT TATGCACCCCT CCAAGCGGTG 120  
GGAGAAGAAT TCGGCCAGC CAGAGTGCAT GTACCTTTT CTCTCTCACA CTTGAAGCAA 180  
ATTAAAAATAG ACNTAGGTNA ATTNTCAGAT AGCCCTGATG GYTATATTGA TGTTTTACAA 240  
GGATTAGGAC AATCCTTGA TCTGACATGG AGAGATATAA TATTACTGCT AAATCAGACG 300  
15 CTAACCTCAA ATGAGAGAAG TGCTGCCATA ACTGGAGCCC GAGAGTTGG CAATCTCTGG 360  
TATCTCAGTC AGGTCAATGA TAGGATGACA ACGGAGGAAA GAGAACGATT CCCCCACAGGG 420  
CAGCAGGCAG TTCCCAGTGT AGCTCCTCAT TGGGACACAG AATCAGAACAA TGGAGATTGG 480  
TGCCGCAGAC ATTAA 495

20

(2) INFORMATION FOR SEQ ID NO: 60:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 2503 base pairs
- (B) TYPE: nucleotide
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

30

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 60:

CCAAGAACCC ACCAATTCCG GANCACATTT TGGCGACCAC GAAGGGACTT TCGCATATCG 60  
CCAAGCGGTG AGACAATAGC CGAGCGGTGA GACCTTCCC AATCGCCAAG CAGTGAGTAC 120  
35 CATCAGACCC CTTTCACTTG CTATTCTGTC CTATCTTCT TTAGAATTGG GGGGCTAAAT 180  
ACCGGGCATC TGTCAGCCAT TTAAAAGTGA CTAGCGGGCC GCCGGACTAA AGACACGGGT 240

GTCAAGCTTT CTGGGAAAGG GCTCTCTAAC AACCCCCAAC TCTTGAGT TGGGACCGTT 300  
 GGTTTGCCTA GAACCAGCTT CCGCTTTCC TGTACTTCTG GGCTGAGCCG TGGGTTGACA 360  
 GTGAAGGAAA GCCATGCATC TCCGGGGTCT CGMCAACATG TTGGTGTGACC CTGCCGCCAT 420  
 GAGTGGAACT CTCAAAAGCA TGTCGCCAA GCGACACTCG CCTATCTATC CTATCTATCC 480  
 5 TGACCCCTG CTCCTGGTC CTAATGCCTG CCAGACAAAC TTCCCTCTCGC CTCTCTTCTC 540  
 TGAAGCTAGA ACCGCTTCTA AAAATTGCTA CCTGGTCTCT GGTGCTTTTC CTARTTCTC 600  
 CTATAAAAGAA TGAWTTCTAG TATTAAACTC CAGGACTCTG TTACCTTCTT TAGGCACCCG 660  
 GGCTCACCAA TCAGAAAGAC ACAGTTTTG CCCAAGGCCG CATCGTAGTG GGGACTACCT 720  
 GGAATTTAG GATCCCTCCT CAGACTAACAA GGCTAACAA AAGTTATTCC TGAAGCTAGG 780  
 10 ATATGGGAG CCTCAGAAAT TGTATCCCTC CTATTCAAT AAGTGAGAAC AAAAGGTGTC 840  
 ACTCTTCCAA CCCTGAAGAT CCCCTCCCTC CCTCAGGGTA TGGCCCTCCA TTTCATTTT 900  
 GTGGCATAAC ATCTTTATAG GATGGGGTAA AGTCCAATA CTAACAGGAG AATGCTTAGG 960  
 ACTCTAACAG GTTTTGAGA ATGCAGTCAGT AAGGGCCACT AAATCTGATT TTTCTCAGTC 1020  
 GGTCCCTCCTT GTGGTCTAGG AGGACAGGCA AGGTTGTGCA GGTTTTCAG AATGCCGTAG 1080  
 15 TAAGGACCCAC TAAATCCGAC CTTCCCTCGGT CCTCCATGTG GTCTGGGAGG AAAACTAGTG 1140  
 TTTCTGCTGC TCGCTCGGTG AGCGCAACTA TTCAAGTCAG CAGGGTCCAG GGACCGTTGC 1200  
 AGGTTCTTGG GCAGGGGTTG TTTCTGCTGC TGCATTGGTG AATGCAACTA TTCTGATCAG 1260  
 CAGGGTCCCA GGACCATTGC AGGTCCCTGG GCAGGGAGAG AAACAAAACA AACCAAAACT 1320  
 GTGGGCGGTT TTGTCTTCA TATGGGAAAC ACTCAGGCAT CAACAGGTTG ACCCTTGAAA 1380  
 20 TGCATCCTAA GCCATTGGGA CCAATTGAC CCACAAACCC TGAAAAAGAG GAGGCTCATT 1440  
 TTTCTGCA CTACGGCTTG GCCCAATAT TCTCTTYTG ATGGGAAAA ATGGCCACCT 1500  
 GAGGGAAGCA CAAATTACAA TAYTATCCTA CAGCYTGATC TTTCTGTAA GAGGGAAGGC 1560  
 AAATGGAGTG AATACCTTAT GTCCAAGCTT TCTTTCTATT GAGGGAGAAT ACACAACAT 1620  
 GCAAAGCTTG CAATTTACAT CCCACAGGAG GACCCCTCAG CTTACCCCCA TATCCTAGCC 1680  
 25 TCCCTATAGC TTCCCTTCCT ATTGATGATA CTCCTCCCT AATCTCCCCT GCCCAGAAGG 1740  
 AAATAAGCAA AGAAATCTCC AAAGGTCCAC AAAACCCCCC GGGCTATCGG TTATGCTCCCT 1800  
 TCAAGYTGTA GGGGGAGGGG AATTGGCCC AACCCGGGTG CATGTCCCTT CTCCCTCTCT 1860  
 GATTTAAAGC AGATCAAGGC AGACCTGGGG AAGTTTCAG ATGATCCTGA TAGGTACATA 1920  
 GATGTCCTAC AGGGTCTAGG GCAAACCTTT GACCTCACTT GGAGAGACGT CATGCTACTG 1980  
 30 TTAGATCAAA CCCTGGCCTT TAATGAAAAG AATGCGGCTT TAGCTGCAGC CTGAGAGTTT 2040  
 GGAGATACCT GGTATCCTAG TCAAGTAAAT GAAAGAATGA CAGCCGAAGA AAGGGACAAC 2100  
 TTCCTTACTG GTCAGCAACC CATCCCCAGT ATGGATCCCC ACTGGGACTT TGACTCAGAT 2160  
 CATGGGGACT GGAGTCGTTA ACATCTGTTG ATCTGTGTTG TGGAAGGACT AAGGAGAATT 2220  
 GGGAAAAAGC CCATGAATTA TTCAATGATA TCCACCATAA CCCAGGGAAA GGAAGAAAAT 2280  
 35 CCTTCTGCCT TCCTCGAGCG GCTACAAGAG GCCTTAAGAA AATATACTCC CCTGTCACCC 2340  
 GAATCACTCG AGGGTCAATT GATTCTAAAA GATAAGTTA TTACCCAATC AGCCACAGAT 2400

ATCAGGAGAA AGCTCCAAA GCAAGCCCTG AGCCTGAACA AAATCTAGAG ACATTATTAA 2460  
ACCTGGCAAC CTTGGTGTTC TATAATAGGG ACCAAGAGGA ACA 2503

## 5 (2) INFORMATION FOR SEQ ID NO: 61:

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 1167 base pairs  
(B) TYPE: nucleotide  
10 (C) STRANDEDNESS: single  
(D) TOPOLOGY: linear

## (ii) MOLECULE TYPE: cDNA

## 15 (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 61:

AAGGAAACTC AGAAAGCAA TACCCATTGTA GTAAAGATGGA CACCAGAAC AGAACAGCT 60.  
TTCCAGGCCA TAAAGAAATC CCTAACCCAA GCCCCAGTGT TAAGCTTGCC AACGGGGCAA 120  
GACTTTCTT TATATGTAC AGAAAAACAG GAATAGCTCT AGGAGTCCTT ACACAGGTCC 180  
20 AAGGGACAAG CTTGCAACCT GTGGCATACC TGAGTAAGGA AACTGATGTA NTGGCAAAGG 240  
GTTGGCCTCA TTGTTTACAG GTAGGGCAGC AGTAGCAGTC TTAGTTCTG AAACAGTTAA 300  
AATAATACAG GGAAGAGATC TTACTGTGTG GACATCTCAT GATGTGAACG GCATAACTCAC 360  
TGCTAAAGAG GACTTGTGGC TGTCAGACAA CCATTTACTT AAATAGCAGG TTCTATTACT 420  
TGAAGTGCCA GTGCTGCGAC TGCACATTG TGCAACTCTT AACCCAGCCA CATTCTTCC 480  
25 AGACAATGAA GAAAAGATAG AACATAACTG TCAACAAGTA ATTGCTAAA CCTATGCTGC 540  
TCGAGGGGAC CTTCTAGAGG TTCCCTTGAC TGATCCGAC CTCAACTTGT ATACTGATGG 600  
AAGTTCTTG GCAGAAAAAG GACTTGAAA AGCGGGGTAT GCAGTGATCA GTGATAATGG 660  
AATACTTGAA AGTAATGCC TCACTCCAGG AACTAGTGCT CACCTGGCAG AACTAATAGC 720  
CCTCACTTGG GCACTAGAAT TAGGAGAAGG AAAAAGGGTA AATATATATT CAGACTCTAA 780  
30 GTATGCTTAC CTAGTCCTCC ATGCCATGC AGCAATATGG AGAGAGAGGG AATTCTAAC 840  
TTCTGAGGGA ACACCTATCA ACCATCAGGG AAGCCATTAG GAGATTATTA TTGGCTGTAC 900  
AGAAACCTAA AGAGGTGGCA GTCTTACACT GCCAGGGTCA TCAGGAAGAA GAGGAAAGGG 960  
AAATAGAAGG CAATGCCAA GCGGATATTG AAGCAAAAAA AGCCGCAAGG CAGGACTCTC 1020  
CATTAGAAAT GCTTATAGAA GGACCCCTAG TATGGGGTAA TCCCCTCTGG GAAACCAAGC 1080  
35 CCCAGTACTC AGCAGGAAAA ATAGAATAGG AAACCTCACA AGGACATACT TTCCTCCCC 1140  
CCAGATGGCT AGCCACTGAG GAAGGAA 1167

## (2) INFORMATION FOR SEQ ID NO: 62:

## 5 (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 78 base pairs
- (B) TYPE: nucleotide
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

10

## (ii) MOLECULE TYPE: cDNA

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 62:

15	TCCAAAGGCA CCAGGGCCCT CAGTGAGGAA CGTATCCAGC CTATACTGGC TTATCCTCAT	60
	CCCCAAAACCC TAAAGCAA	78

## (2) INFORMATION FOR SEQ ID NO: 63

20

## (i) SEQUENCE CHARACTERISTICS :

- (A) LENGTH : 26 amino acids
- (B) TYPE : amino acid

25 (ii) MOLECULE TYPE : peptide

## (xi) SEQUENCE DESCRIPTION : SEQ ID NO: 63

Ser Lys Gly Thr Arg Ala Leu Ser Glu Glu Arg Ile Gln Pro Ile Leu				
30	1	5	10	15
Ala Tyr Pro His Pro Lys Thr Leu Lys Gln				
	20		25	

35 (2) INFORMATION FOR SEQ ID NO: 64:

159

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 28 base pairs
- (B) TYPE: nucleotide
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

5

## (ii) MOLECULE TYPE: cDNA

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 64:

10

AAATGTCTGC GGCACCAATC TCCATGTT

28

## (2) INFORMATION FOR SEQ ID NO: 65:

15

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 30 base pairs
- (B) TYPE: nucleotide
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

20

## (ii) MOLECULE TYPE: cDNA

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 65:

25

AAGGGGCATG GACGAGGTGG TGGCTTATTT

30

## (2) INFORMATION FOR SEQ ID NO: 66:

30

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 21 base pairs
- (B) TYPE: nucleotide
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

35

160

(ii) MOLECULE TYPE: cDNA

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 66:

GGAGAAAGAGC AGCATAAGTG G 21

5

(2) INFORMATION FOR SEQ ID NO: 67:

(i) SEQUENCE CHARACTERISTICS:

- 10 (A) LENGTH: 25 base pairs  
(B) TYPE: nucleotide  
(C) STRANDEDNESS: single  
(D) TOPOLOGY: linear

15 (ii) MOLECULE TYPE: cDNA

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 67:

GTGCTGATTG GTGTATTTAC AATCC 25

20

(2) INFORMATION FOR SEQ ID NO: 68:

(i) SEQUENCE CHARACTERISTICS:

- 25 (A) LENGTH: 34 base pairs  
(B) TYPE: nucleotide  
(C) STRANDEDNESS: single  
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

30

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 68:

GACTCGCTGC AGATCGATTT TTTTTTTTTT TTTT 34

35 (2) INFORMATION FOR SEQ ID NO: 69:

161

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 30 base pairs
- (B) TYPE: nucleotide
- (C) STRANDEDNESS: single
- 5 (D) TOPOLOGY: linear

## (ii) MOLECULE TYPE: cDNA

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 69:

10

GCCATCAAGC CACCCAAGAA CTCTTAAC TT

30

## (2) INFORMATION FOR SEQ ID NO: 70:

15

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 30 base pairs
- (B) TYPE: nucleotide
- (C) STRANDEDNESS: single
- 20 (D) TOPOLOGY: linear

## (ii) MOLECULE TYPE: cDNA

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 70:

25

CCAATAGCCA GACCATTATA TACACTAATT

30

## (2) INFORMATION FOR SEQ ID NO: 71:

30

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 23 base pairs
- (B) TYPE: nucleotide
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

35

## (ii) MOLECULE TYPE: cDNA

162

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 71:  
GCCATAACTG CAACCCAAGA GTT

23

5

(2) INFORMATION FOR SEQ ID NO: 72:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 23 base pairs
- 10 (B) TYPE: nucleotide
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

15

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 72:

GGACGAGGTG GTGGCTTATT TCT

23

20

(2) INFORMATION FOR SEQ ID NO: 73:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 25 base pairs
- 25 (B) TYPE: nucleotide
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

30

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 73:

AACTTGCCTG CTAGAAGGAC TAAGG

25

35

(2) INFORMATION FOR SEQ ID NO: 74:

163

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 24 base pairs
- (B) TYPE: nucleotide
- 5 (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

## (ii) MOLECULE TYPE: cDNA

10 (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 74:

AACTTTCCC TTTTCCAGAT CCTC

24

15 (2) INFORMATION FOR SEQ ID NO: 75:

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 22 base pairs
- (B) TYPE: nucleotide
- 20 (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

## (ii) MOLECULE TYPE: cDNA

25 (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 75:

GCATACCAAGG CAAGTGGACA TT

22

30 (2) INFORMATION FOR SEQ ID NO: 76:

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 25 base pairs
- (B) TYPE: nucleotide
- 35 (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

164

(ii) MOLECULE TYPE: cDNA

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 76:

5

CTGTCCGTTG GGTTTCCTTA CTCCT

25

(2) INFORMATION FOR SEQ ID NO: 77:

10

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 24 base pairs
- (B) TYPE: nucleotide
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

15

(ii) MOLECULE TYPE: cDNA

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 77:

20

GAGGCTCTGG AAAAGGGAAA AGTT

24

(2) INFORMATION FOR SEQ ID NO: 78:

25

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 25 base pairs
- (B) TYPE: nucleotide
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

30

(ii) MOLECULE TYPE: cDNA

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 78:

35

CTGTCCGTTG GGTTTCCTTA CTCCT

25

(2) INFORMATION FOR SEQ ID NO: 79:

5 (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 25 base pairs
- (B) TYPE: nucleotide
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

10

(ii) MOLECULE TYPE: cDNA

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 79:

15 AGGAGTAAGG AAACCCAACG GACAG

25

(2) INFORMATION FOR SEQ ID NO: 80:

(i) SEQUENCE CHARACTERISTICS:

- 20
- (A) LENGTH: 25 base pairs
  - (B) TYPE: nucleotide
  - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear

25 (ii) MOLECULE TYPE: cDNA

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 80:

TGTATATAAT GGTCTGGCTA TTGGG

25

30

(2) INFORMATION FOR SEQ ID NO: 81:

(i) SEQUENCE CHARACTERISTICS:

- 35
- (A) LENGTH: 25 base pairs
  - (B) TYPE: nucleotide

166

- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

5

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 81:

AGGAGTAAGG AAACCCAACG GACAG

25

10

(2) INFORMATION FOR SEQ ID NO: 82:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 26 base pairs
- (B) TYPE: nucleotide
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

20

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 82:

TTCGGCAGAA ACCTGTTATG CCAAGG

26

25

(2) INFORMATION FOR SEQ ID NO: 83:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 22 base pairs
- (B) TYPE: nucleotide
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

35

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 83:

167

CTCGATTCT TGCTGGCCT TA

22

5 (2) INFORMATION FOR SEQ ID NO: 84:

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 20 base pairs
- (B) TYPE: nucleotide
- 10 (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

## (ii) MOLECULE TYPE: cDNA

15 (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 84:

GTTGATTCCC TCCTCAAGCA

20

20 (2) INFORMATION FOR SEQ ID NO: 85:

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 20 base pairs
- (B) TYPE: nucleotide
- 25 (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

## (ii) MOLECULE TYPE: cDNA

30 (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 85:

CTCTACCAAT CAGCATGTGG

20

35 (2) INFORMATION FOR SEQ ID NO: 86:

168

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 19 base pairs
- (B) TYPE: nucleotide
- (C) STRANDEDNESS: single
- 5 (D) TOPOLOGY: linear

## (ii) MOLECULE TYPE: cDNA

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 86:

10

TGTTCCCTCTT GGTCCCTAT

19

## (2) INFORMATION FOR SEQ ID NO: 87:

15

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 433 aminoacids
- (B) TYPE: aminoacid

20

## (ii) MOLECULE TYPE: peptide

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 87:

Met Ala Thr Ala Thr Gly Thr Gly Ile Ala Gly Leu Ser Thr Ser Leu				
1	5	10	15	
Ser Tyr Tyr His Thr Leu Ser Lys Asn Phe Ser Asp Ser Leu Gln Glu				
20	25	30		
Ile Met Lys Ser Ile Leu Thr Leu Gln Ser Gln Leu Asp Ser Leu Ala				
35	40	45		
Ala Met Thr Leu Gln Asn Arg Arg Gly Pro His Leu Leu Thr Ala Glu				
30	50	55	60	
Lys Gly Gly Leu Cys Thr Phe Leu Gly Glu Glu Cys Cys Phe Tyr Thr				
65	70	75	80	
Asn Gln Ser Gly Ile Val Arg Asp Ala Thr Trp His Leu Gln Glu Arg				
85	90	95		
35 Ala Ser Asp Ile Arg Gln Cys Leu Ser Asn Ser Tyr Thr Asn Leu Trp				
100	105	110		

169

Ser Trp Ala Thr Trp Leu Leu Pro Phe Leu Gly Pro Met Ala Ala Ile  
115 120 125  
Leu Leu Leu Leu Thr Phe Gly Pro Cys Ile Phe Lys Leu Leu Val Lys  
130 135 140  
5 Phe Val Ser Ser Arg Ile Glu Ala Ile Lys Leu Gln Met Val Leu Gln  
145 150 155 160  
Met Glu Pro Gln Met Ser Ser Thr Asn Asn Phe Tyr Gln Gly Pro Leu  
165 170 175  
Glu Arg Ser Thr Gly Thr Ser Thr Ser Leu Glu Ile Pro Leu Trp Lys  
10 180 185 190  
Thr Leu Gln Leu Gln Gly Pro Phe Phe Ala Pro Ile Gln Gln Glu Val  
195 200 205  
Ala Arg Ala Val Ile Gly Gln Ile Pro Asn Ser Ser Trp Gly Val Leu  
210 215 220  
15 Phe Arg Gly Gly Ile Glu Glu Val Thr Ala Cys Trp Gln Pro His Ser  
225 230 235 240  
Pro Arg Trp Xaa Ser Val Pro Pro Gln Pro Trp Cys Pro Leu Trp Pro  
245 250 255  
Cys Leu Arg Ser Pro Ser Ala Cys His Cys Thr Val Gly Ala Ser Phe  
20 260 265 270  
Trp Ala Gly Gln Gly Arg Ser Gln Leu Pro Gln Leu Ala Gly Arg Tyr  
275 280 285  
Gly Gly Arg Asp Ala Gly Gly Asn Gln Gly Cys Ala Trp Arg Leu Arg  
290 295 300  
25 Ala Ser Met Ser Ser Arg Trp Ala Trp Ala Arg Arg Ala Pro His Ser  
305 310 315 320  
Gly Ser Glu Gly Leu Ser Thr Trp Ala Arg Gln Met Leu Cys Ser Thr  
325 330 335  
Ser Ser Leu Gly Leu Ser Cys Leu Pro Arg Gly Ala Gly Leu Arg Glu  
30 340 345 350  
Xaa Ala Ala Cys Pro Cys Leu Ser Pro Pro Pro Arg Arg Gly Phe Leu  
355 360 365  
His Ser Pro Ser Phe Pro Asp Lys His His Pro Leu Ser Thr Val Pro  
370 375 380  
35 Ser Pro Ile Asn His Pro Arg Val Glu Glu Cys Gly His Thr Ala Arg  
385 390 395 400

170

Asp Trp Gln Ala Val Pro Leu Ala Ala Leu Val Arg Asp Pro Leu Arg  
405 410 415  
Glu Ala Ser Trp Ala Pro Glu Ser Gly Gly Asp Leu Glu Asn Leu Tyr  
420 425 430  
5 Val  
433

## (2) INFORMATION FOR SEQ ID NO: 88:

10

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 693 base pairs
- (B) TYPE: nucleotide
- (C) STRANDEDNESS: single
- 15 (D) TOPOLOGY: linear

## (ii) MOLECULE TYPE: cDNA

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 88:

20

CTTCCCCAAC TAATAAGGAC CCCCTTTCA ACCAAACAG TCCAAAGGA CATAGACAAA 60  
GGAGTAAACA ATGAACCAA GAGTCCAAT ATTCCCTGGT TATGCACCCCT CCAAGCGGTG 120  
GGAGAAGAAT TCGGCCAGC CAGAGTCAT GTACCTTTT CTCTCTCACA CTTGAAGCAA 180  
ATTAAAATAG ACNTAGGTNA ATTNTCAGAT AGCCCTGATG GYTATATTGA TGTTTACAA 240  
25 GGATTAGGAC AATCCTTGTA TCTGACATGG AGAGATATAA TATTACTGCT AAATCAGACG 300  
CTAACCTCAA ATGAGAGAAAG TGCTGCCATA ACTGGAGCCC GAGAGTTGG CAATCTCTGG 360  
TATCTCAGTC AGGTCAATGA TAGGATGACA ACGGAGGAAA GAGAACGATT CCCCCACAGGG 420  
CAGCAGGCAG TTCCCAAGTGT AGCTCCTCAT TGGGACACAG AACAGAACAA TGGAGATTGG 480  
TGCCGCAGAC ATTTACTAAC TTGCGTGCTA GAAGGACTAA GGAAAACTAG GAAGACTATG 540  
30 AATTATTCAA TGATGTCCAC TATAACACAG GGGAAAGGAA GAAAATCCTA CTGCCTTTCT 600  
GGAGAGACTA AGGGAGGCAT TGAGGAAGCA TACCAAGCAA GTGGACATTG GAGGCTCTGG 660  
AAAAAGGGAAA AGTTGGGCAA ATTGAATGCC TAA 693

35 (2) INFORMATION FOR SEQ ID NO: 89:

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 1577 base pairs
- (B) TYPE: nucleotide
- (C) STRANDEDNESS: single
- 5 (D) TOPOLOGY: linear

## (ii) MOLECULE TYPE: cDNA

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 89:

10 AACTTGCCTG CTAGAAGGAC TAAGGAAAAC TAGGAAGACT ATGAATTATT CAATGATGTC 60  
CACTATAACA CAGGGGAAAG GAAGAAAATC CTACTGCCCT TCTGGAGAGA CTAAGGGAGG 120  
CATTGAGGAA GCATACCAGG CAAGTGGACA TTGGAGGCTC TGAAAAGGG AAAAGTTGGG 180  
CAAATTGAAT GCCTAATAGG GCTTGCTTCC AGTGCAGTCT ACAAGGACGC TTTAGAAAAG 240  
15 ATTGTCCAAG TAGAAATAAG CCGCCCCCTCG TCCATGCCCT TTATGTCAAG GGAATCACTG 300  
GAAGGCCTAC TGCCCCAGGG GACGAAGGTC CTCTGAGTCA GAAGCCACTA ACCTGATGAT 360  
CCAGCAGCAG GACTGAGGGT GCCCGGGGCA AGTGCCAGCC CATGCCATCA CCCTCAGAGC 420  
CCC GG GTATG TTTGACCATT GAGAGCCAGG AAGTTAACTG TCTCCTGGAC ACTGGCGCAG 480  
CCTTCTCAGT CTTACTTTCC TGTC CAGAC AATTGTCTC CAGATCTGTC ACTATCCGAG 540  
20 GGGTCCTAAG ACAGCCAGTC ACTACATACT TCTCTCAGCC ACTAAGTTGT GACTGGGAA 600  
CTTTACTCTT TTCACATGCT TTTCTAATTA TGCCTGAAAG CCCCACCTCC TTGTTAGGGA 660  
GAGACATTTT AGCAAAGCA GGGGCCATTA TACACCTGAA CATAGGAAA GGAATACCCA 720  
TTTGCTGTCC CCTGCTTGAG GAAGGAATTAA ATCCTGAAGT CTGGCAATA GAAGGACAAT 780  
ATGGACAAGC AAAGAATGCC CGTCCTGTTC AAGTTAAACT AAAGGATTCT GCCTCCTTTC 840  
25 CCTACCAAAG GAAGTACCCCT CTTAGACCCG AGGCCCTACA AGGACTCAAA AGATTGTTAA 900  
GGACCTAAAA GCCCAAGGCC TAGTAAAACC ATGCAGTAGC CCCTGCAATA CTCCAATTAA 960  
AGGAGTAAGG AAACCCAACG GACAGTGGAG GTTAGTGCAGA GATCTCAGGA TTATTAATGA 1020  
GGCTGTTTT CCTCTATACC CAGCTGTATC TAGCCCTTAT ACTCTGCTTT CCCTAATACC 1080  
AGAGGAAGCA GAGTAGTTA CAGTCCTGGA CCTTAAGGAT GCCTCTTCT GCATCCCTGT 1140  
30 ACATCCTGAT TCTCAATTCT TGTTGTCCT TGAAGATCCT TTGAACCCAA TGTCTCAATT 1200  
CACCTGGACT GTTTACCC AGGGGTTCCG GGATAGCCCA CATCTATTG GCCAGGCATT 1260  
AGCCCAAGAC TTGAGCCAAT TCTCATACCT GGACATCTG TCCTTCGGTA TGGGATGATT 1320  
TAATTTAGC CACCCGTTCA GAAACCTTGT GCCATCAAGC CACCCAAAGCG TTCTTAAATT 1380  
TCCTCACTCC GTGTGGCTAC AAGGTTCCA AACCAAAGGC TCAGCTCTGC TCACAGCAGG 1440  
35 TTAAATACTT AGGGTTAAAA TTATCCAAAG GCACCCAGGGC CCTCTGTGAG GAATGTATCC 1500  
AACCTGTACT GGCTTATCTT CATCCCAAAA CCCTAAAGCA ACTAAGAAGG TCCTTGGCAT 1560

172

AACAGGTTTC TGCCGAA

1577

## (2) INFORMATION FOR SEQ ID NO: 90:

5

## (i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 182 amino acids

(B) TYPE: amino acid

10 (ii) MOLECULE TYPE: peptide

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 90:

Ser Ser Ser Arg Thr Glu Gly Ala Arg Gly Lys Cys Gln Pro Met Pro  
15 1 5 10 15

Ser Pro Ser Glu Pro Arg Val Cys Leu Thr Ile Glu Ser Gln Glu Val  
20 20 25 30

Asn Cys Leu Leu Asp Thr Gly Ala Ala Phe Ser Val Leu Leu Ser Cys  
35 35 40 45

20 Pro Arg Gln Leu Ser Ser Arg Ser Val Thr Ile Arg Gly Val Leu Arg  
50 50 55 60

Gln Pro Val Thr Thr Tyr Phe Ser Gln Pro Leu Ser Cys Asp Trp Gly  
65 65 70 75 80

Thr Leu Leu Phe Ser His Ala Phe Leu Ile Met Pro Glu Ser Pro Thr  
25 85 90 95

Pro Leu Leu Gly Arg Asp Ile Leu Ala Lys Ala Gly Ala Ile Ile His  
100 100 105 110

Leu Asn Ile Gly Lys Gly Ile Pro Ile Cys Cys Pro Leu Leu Glu Glu  
115 115 120 125

30 Gly Ile Asn Pro Glu Val Trp Ala Ile Glu Gly Gln Tyr Gly Gln Ala  
130 130 135 140

Lys Asn Ala Arg Pro Val Gln Val Lys Leu Lys Asp Ser Ala Ser Phe  
145 145 150 155 160

Pro Tyr Gln Arg Lys Tyr Pro Leu Arg Pro Glu Ala Leu Gln Gly Leu  
35 165 170 175

Lys Arg Leu Leu Arg Thr

173

180

## (2) INFORMATION FOR SEQ ID NO: 91:

5

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 36 base pairs
- (B) TYPE: nucleotide
- (C) STRANDEDNESS: single
- 10 (D) TOPOLOGY: linear

## (ii) MOLECULE TYPE: cDNA

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 91:

15

AGATCTGCAG AATTCGATAT CACCCCCCCC CCCCCC

36

## (2) INFORMATION FOR SEQ ID NO: 92:

20

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 22 base pairs
- (B) TYPE: nucleotide
- (C) STRANDEDNESS: single
- 25 (D) TOPOLOGY: linear

## (ii) MOLECULE TYPE: cDNA

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 92:

30

AGATCTGCAG AATTCGATAT CA

22

## (2) INFORMATION FOR SEQ ID NO: 93:

## (i) SEQUENCE CHARACTERISTICS:

35

- (A) LENGTH: 2304 base pairs
- (B) TYPE: nucleotide

(C) STRANDEDNESS: single

(D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 93:

5	TCCAGCAGCA GGACTGAGGG TGCCCGGGGC AAGTGCCAGC CCATGCCATC ACCCTCAGAG CCCCGGGTAT GTTGACCAT TGAGAGCCAG GAAGTTAACT GTCTCCTGGA CACTGGCGCA GCCTCTCAG TCTTACTTTC CTGTCCCAGA CAATTGTCCT CCAGATCTGT CACTATCCGA GGGGTCCCTAG GACAGCCAGT CACTACATAC TTCTCTCAGC CACTAAGTTG TGACTGGGGA ACTTTACTCT	50 100 150 200 250
10	TTTCACATGC TTTTCTAATT ATGCCTGAAA GCCCCACTCC CTTGTTAGGG AGAGACATTT TAGCAAAAGC AGGGGCCATT ATACACCTGA ACATAGGAAA AGGAATACCC ATTTGCTGTC CCCTGCTTGA GGAAGGAATT AATCCTGAAG TCTGGGCAAT AGAAGGACAA TATGGACAAG CAAAGAATGC CCGTCCTGTT CAAGTTAACAC TAAAGGATTC TGCCCTCCTT CCCTACCAAA GGAAGTACCC	300 350 400 450 500
15	TCTTAGACCC GAGGCCCTAC AAGGANCTCA AAAGATTGTT AAGGACCTAA AAGCCAAGG CCTAGTAAAA CCATGCAGTA GCCCCTGCAA TACTCCAATT TTAGGAGTAA GGAAACCCAA CGGACAGTGG AGGTTAGTGC AAGATCTCAG GATTATTAAT GAGGCTGTTT TTCCTCTATA CCCAGCTGTA TCTAGCCCTT ATACTCTGCT TTCCCTAATA CCAGAGGAAG CAGAGTGGTT TACAGTCCTG	550 600 650 700 750
20	GACCTTAAGG ATGCCTTTT CTGCATCCCT GTACGTCCCTG ACTCTCAATT CTTGTGTTGCC TTTGAAGATC CTTTGAACCC AACGTCTCAA CTCACCTGGA CTGTTTACCC CCAAGGGTTC AGGGATAGCC CCCATCTATT TGGCCAGGCA TTAGCCCAAG ACTTGAGTCA ATTCTCATAC CTGGACACTC TTGTCCTTCA GTACGTGGAT GATTTACTTT TAGTCGCCCG TTCAGAAACC TTGTGCCATC	800 850 900 950 1000
25	AAGCCACCCA AGAACTCTTA ACTTTCTCA CTACCTGTGG CTACAAGGTT TCCAAACCAA AGGCTCGGCT CTGCTCACAG GAGATTAGAT ACTTAGGGCT AAAATTATCC AAAGGCACCA GGGCCCTCAG TGAGGAACGT ATCCAGCCTA TACTGGCTTA TCCTCATCCC AAAACCTAA AGCAACTAAG AGGGTCCTT GGCATAACAG GTTTCTGCCG AAAACAGATT CCCAGGTACA CCCCAATAGC	1050 1100 1150 1200 1250
30	CAGACCATTA TATACACTAA TTAGGGAAAC TCAGAAAGCC AATACCTATT TAGTAAGATG GACACCTACA GAAGTGGCTT TCCAGGCCCT AAAGAAGGCC CTAACCCAAG CCCCAGTGTGTT CAGCTTGCCA ACAGGGCAAG ATTTTCTTT ATATGCCACA GAAAAAACAG GAATAGCTCT AGGAGTCCTT ACCGAGGTCT CAGGGATGAG CTTGCAACCC GTGGTATACC TGAGTAAGGA AATTGATGTA	1300 1350 1400 1450 1500
35	GTGGCAAAGG GTTGGCCTCA TTGTTATGG GTAATGGCGG CAGTAGCAGT CTTAGTATCT GAAGCAGTTA AAATAATACA GGGAAAGAGAT CTTACTGTGT	1550 1600

	GGACATCTCA TGATGTGAAC GGCATACTCA CTGCTAAAGG AGACTTGTGG	1650
	TTGTCAGACA ACCATTTACT TAATTATCAG GCTCTATTAC TTGAAGAGCC	1700
	AGTGCTGAGA CTGCGCACTT GTGCAACTCT TAAACCCGCC ACATTTCTTC	1750
	CAGACAATGA AGAAAAGATA GAACATAACT GTCAACAAAGT AATTGCTCAA	1800
5	ACCTATGCTG CTCGAGGGGA CCTTCTAGAG GTTCCCTTGA CTGATCCCGA	1850
	CCTCAACTTG TATACTGATG GAAGTTCCCTT GGCAAGAAAAA GGACTTCGAA	1900
	AAGCGGGTA TGCACTGATC AGTGATAATG GAATACCTGA AAGTAATCGC	1950
	CTCACTCCAG GAACTAGTGC TCACCTGGCA GAACTAATAG CCCTCACTTG	2000
	GGCACTAGAA TTAGGAGAAG GAAAAAGGGT AAATATATAT TCAGACTCTA	2050
10	AGTATGCTTA CCTAGTCCTC CATGCCCATG CAGCAATATG GAGAGAGAGG	2100
	GAATTCCCTAA CTTCTGAGGG AACACCTATC AACCATCAGG AAGCCATTAG	2150
	GAGATTATTA TTGGCTGTAC AGAAACCTAA AGAGGTGGCA GTCTTACACT	2200
	GCCAGGGTCA TCAGGAAGAA GAGGAAAGGG AAATAGAAGG CAATGCCAA	2250
	CGGGATATTG AAGCAAAAAA AGCCGCAAGG CAGGACTCTC CATTAGAAAT	2300
15	GCTT	2304

## (2) INFORMATION FOR SEQ ID NO: 94:

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 2364 base pairs
- 20 (B) TYPE: nucleotide
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 94:

25	ATGATCCAGC AGCAGGACNG AGGGTCCCCG GGGCAAGCGC CAGCCCATGC	50
	CATCACCCCTC ACAGAGCCCC AGGTATGCTT GACCATTGAG GGTCAAGAAGG	100
	GTNACTGTCT CCTGGACACT GCGGNGCCT TCTCAGTCTT ACTTTCTGT	150
	CCTGGACAAAC TGTCCCTCCAG ATCTGTCACT GTCCGAGGGG TCCTAGGACA	200
	GCCAGTCACT AGATACTTCT CCCAGCCACT AAGTTGTGAC TGGGGAACTT	250
30	TACTCTTCCC ACATGCTTT CTAATTATGC CTGAAAGCCC CACTCTCTTG	300
	TTGGGGAGAG ACATTCTAGC AAAAGCAGGG GCCATTATAC ATGTGAATAT	350
	AGGAGAAGGA ACAACTGTTT GTTGTCCCCT GCTTGAGGAA GGAATTAATC	400
	CTGAAGTCCG GGCAACAGAA GGACAATATG GACAAGCAAA GAATGCCCGT	450
	CCTGTTCAAG TTAAACTAAA GGATTCCACC TCCTTCCCT ACCAAAGGCA	500
35	GTACCCCTC AGACCCGAGA CCCAACAAAGA ACTCCAAAAG ATTGTAAAGG	550
	ACCTAAAAGC CCAAGGCCTA GTAAAACCAA GCAATAGCCC TTGCAAGACT	600

	CCAATTTAG GAGTAAGGAA ACCCAACGGA CAGTGGAGGT TAGTGCAAGA	650
	ACTCAGGATT ATCAATGAGG CTGTTGTTCC TCTATAACCA GCTGTACCTA	700
	ACCCTTATAC AGTGCTTCC CAAATACCGAG AGGAAGCAGA GTGGTTTACA	750
	GTCCTGGACC TTAAGGATGC CTTTTCTGC ATCCCTGTAC GTCCTGACTC	800
5	TCAATTCTTG TTTGCCTTG AAGATCCTTT GAACCCAACG TCTCAACTCA	850
	CCTGGACTGT TTTACCCCCAA GGTTCAAGGG ATAGCCCCCA TCTATTGGC	900
	CAGGCATTAG CCCAAGACTT GAGTCAATT TCATACCTGG ACACTCTTGT	950
	CCTTCAGTAC ATGGATGATT TACTTTAGT CGCCCGTTCA GAAACCTTGT	1000
	GCCATCAAGC CACCCAAAGAA CTCTTAACCT TCCTCACTAC CTGTGGCTAC	1050
10	AAGGTTTCCA AACCAAAGGC TCGGCTCTGC TCACAGGGAGA TTAGATACTN	1100
	AGGGCTAAAA TTATCCAAAG GCACCAGGGC CCTCAGTGAG GAACGTATCC	1150
	AGCCTATACT GGCTTATCCT CATCCCAAAA CCCTAAAGCA ACTAAGAGGG	1200
	TTCCCTGGCA TAACAGGTTT CTGCCGAAAA CAGATTCCCA GGTACASCCC	1250
	AATAGCCAGA CCATTATATA CACTAATTAN GGAAACTCAG AAAGCCAATA	1300
15	CCTATTAGT AAGATGGACA CCTACAGAAG TGGCTTCCA GGCCCTAAAG	1350
	AAGGCCCTAA CCCAAGCCCC AGTGTTCAGC TTGCCAACAG GGCAAGATT	1400
	TTCTTATAT GCCACAGAAA AAACAGGAAT AGCTCTAGGA GTCCCTACGC	1450
	AGGTCTCAGG GATGAGCTTG CAACCCGTGG TATACCTGAG TAAGGAAATT	1500
	GATGTAGTGG CAAAGGGTTG GCCTCATNGT TTATGGTAA TGGNGGCAGT	1550
20	AGCAGTCTNA GTATCTGAAG CAGTTAAAAT AATACAGGGAGAGATCTTN	1600
	CTGTGTGGAC ATCTCATGAT GTGAACGGCA TACTSRCTGC TAAAGGAGAC	1650
	TTGTGGTTGT CAGACAACCA TTTACTTAAN TAYCAGGCYY TATTACTTGA	1700
	AGAGCCAGTG CTGNNGACTGC GCACCTGTCC AACTCTAAA CCCAAACTTA	1750
	TGCTGCCAG AAGGATCTT NTAGAGGTCC CCTTAGCCAA CCCTGACCTC	1800
25	AACTATATAT ATACTGATGG AAGTCGTTT GTAGAAAAGG GATTACAAAG	1850
	GGNAGGATAT NCCATAGGTG TTAGTGATAA AGCAGTACTT GAAAGTAAGC	1900
	CTCTTCCCCC CCAGGGACCA GCGCCCCCGT TAGCAGAACT AGTGGCACTG	1950
	ACCCCGCGAG CCTTAGAAGT TTGAAAGGG AGGAGGATAA ATGTGTATAC	2000
	AGATAGCAAG TATGCTTATC TAATCCGAAA TGCCCATGTT GCAATATGGA	2050
30	AAGAAAGGGA GTTCCTAACCC TCTGGGGAA CCCCCATTAA ATACCACAAAG	2100
	TTAACATGG AGTTATTGCA CACAGTGCAA AAACTCAAGG AGGTGGAAGT	2150
	CTTACACTGC CAAAGCCATC AGAAAAGGGAA AAGAGGGAA GAGCAGCATA	2200
	AGTGGCTACA GAGGCAAGGA AAGACTAGCA GAAAGGAAAG AGAGAAAGAG	2250
	ACAGAAAGTC AGAGAGAGAG AGAGGAAGAG ACAGAGCACA AAGAGGGAGT	2300
35	CAGAGAGAGA GAGAGACAGA GAGTCAGAGA GAAGGAAAGA GAGAGAGGAA	2350
	GAGACAAAGA ATGAH	2365

## (2) INFORMATION FOR SEQ ID NO: 95:

## (i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 768 amino acids

5 (B) TYPE: peptide

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 95:

	SSSRTEGARG KCQPMPS PSE PRVCLTIESQ EVNCLLDTGA AFSVLLSCPR	50
	QLSSRSVTIR GVLGQPVTYY FSQPLSCDWG TLLFSHAF LI MPESPTPLL G	100
10	RDILAKAGAI IHLNIGKGIP ICCPLLEEGI NPEVWAIEGQ YGQAKNARPV	150
	QVKLKDSASF PYQRKYPLRP EALQGXQKIV KDLKAQGLVK PCSSPCNTPI	200
	LGVRKPNGQW RLVQDLRIIN EAVFPLYP AV SSPYTLLSLI PEEAEWFTV L	250
	DLKDAFFCIP VRPDSQFLFA FEDPLNPTS Q LTWTVLPQGF RDSPHLFGQ A	300
	LAQDLSQFSY LD TLV LQYVD DLLLVARSET LCHQATQELL TFLTTCGYKV	350
15	SKPKARLCSQ EIRYLGLKLS KGTRALSEER IQPILAYPHP KTLKQLRGFL	400
	GITGFCRKQI PRYTPIARPL YTLIRETQKA NTYLVWRWTPT EVAFQALKKA	450
	L TQAPVFSLP TGQDFSLYAT EKTGIALGVL TQVSGMSLQP VVYLSKEIDV	500
	VAKGWP HCLW VMAAVAVLVS EAVKIIQGRD LTVWTSHDV N GILTAKGDLW	550
	LSDNHLLNYQ ALLLEEPVLR LRTCATLKPA TFLPDNEEKI EHNCQQVIAQ	600
20	TYAARGDLLE VPLTDPDLNL YT DGS SLAEK GLRKAGYAVI SDNGILESNR	650
	LTPGTS AHLA ELIALTWALE LGEGKRVNIY SDSKYAYLVL HAHA AIWRER	700
	EFLTSEGTP I NHQE AIRR LL LAVQKPKEVA VLHCQGHQEE EEREIEGNR Q	750
	ADIEAKKAAR QDSPLEML	768

## 25 (2) INFORMATION FOR SEQ ID NO: 96:

## (i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 114 amino acids

(B) TYPE: peptide

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 96:

30	SSSRTEGARG KCQPMPS PSE PRVCLTIESQ EVNCLLDTGA AFSVLLSCPR	50
	QLSSRSVTIR GVLGQPVTYY FSQPLSCDWG TLLFSHAF LI MPESPTPLL G	100
	RDILAKAGAI IHLN	114

## 35 (2) INFORMATION FOR SEQ ID NO: 97:

## (i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: amino acids

(B) TYPE: peptide

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 97:

5 IGKGIPICCPPLLEEGINPEVWAIEGQYQAKNARPV  
QVKLKDSASFPYQRKYPLRPEALQGXQKIVKDLKAQGLVVKPCSSPCNTPI  
LGVRKPNGQWRLVQDLRIINEAVFPLYPAVSSPYTLLSLIPEEAEWFTVL  
DLKDAFFCIPVRPDSQFLFAFEDPLNPTSSQLTVLQPQFRDSPHLFGQA  
LAQDLSQFSYLDTLVLQYVDDLLLVARSETLCHQATQELLTFLTCGYKV  
10 SKPKARLCSQEIRYGLKLSKGTRALSEERIQPILAYPHPKTLKQLRGFL  
GITGFCKRKQIPRYTPARIAPLYTLIRETQKANTYLVRWTPTEVAFQALKKA  
LTQAPVFSLPTGQDFSLYATEKTGIALGVLTQVSGMSLQPVVYLSKEIDV  
VAKGWPHCLWVMAAVAVLVSEAVKIIQGRDLTVWTSHDVNGILTAKGDLW  
LSDNHLLNYQALLLEEPVRLRRTCATLKPATFLPDNEEKIEHNCQQVIAQ  
15 TYAARGDLLEVPLTDPLNLYTDGSSLAEKGLRKAGYAVISDNGILESNR  
LTPGTS AHLAE LIALT WAEL GEG KRV NI YSDSKYAYLVL HAHAAI WRER  
EFLTSEGTPINHQEAIRRLLLAVQKPKEVAVLHCQGHQEEEEREIEGNRQ  
ADIEAKKAARQDSPLEML

20

(2) INFORMATION FOR SEQ ID NO: 98:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: amino acids

(B) TYPE: peptide

25 (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 98:

LYTDGSSLAEKGLRKAGYAVISDNGILESNR  
LTPGTS AHLAE LIALT WAEL GEG KRV NI YSDSKYAYLVL HAHAAI WRER  
EFLTSEGTPINHQEAIRRLLLAVQKPKEVAVLHCQGHQEEEEREIEGNRQ

30 ADIEAKKAARQDSPLEML

(2) INFORMATION FOR SEQ ID NO: 99

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 23 base pairs

(B) TYPE: nucleotide

(C) STRANDEDNESS: single

35

179

(D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 99:

AGGAGTAAGG AAACCCAACG GAC

23

5 (2) INFORMATION FOR SEQ ID NO: 100

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 19 base pairs

(B) TYPE: nucleotide

(C) STRANDEDNESS: single

10 (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 100:

TAAGAGTTGC ACAAGTGC

19

(2) INFORMATION FOR SEQ ID NO: 101

15 (i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 21 base pairs

(B) TYPE: nucleotide

(C) STRANDEDNESS: single

(D) TOPOLOGY: linear

20 (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 101:

TCAGGGATAG CCCCCATCTA T

21

(2) INFORMATION FOR SEQ ID NO: 102

(i) SEQUENCE CHARACTERISTICS:

25 (A) LENGTH: 24 base pairs

(B) TYPE: nucleotide

(C) STRANDEDNESS: single

(D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 102:

30 AACCCATTGCA CACTACATCA ATTT

24

(2) INFORMATION FOR SEQ ID NO: 103

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 18 base pairs

35 (B) TYPE: nucleotide

(C) STRANDEDNESS: single

180

(D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 103:

AGCAGCAGGA CTGAGGGT

18

5 (2) INFORMATION FOR SEQ ID NO: 104

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 25 base pairs

(B) TYPE: nucleotide

(C) STRANDEDNESS: single

10 (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 104:

CTGTCCGTTG GGTTTCCTTA CTCCT

25

(2) INFORMATION FOR SEQ ID NO: 105

15 (i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 24 base pairs

(B) TYPE: nucleotide

(C) STRANDEDNESS: single

(D) TOPOLOGY: linear

20 (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 105:

GACAGCAAAT GGGTATTCTT TTCC

24

(2) INFORMATION FOR SEQ ID NO: 106

(i) SEQUENCE CHARACTERISTICS:

25 (A) LENGTH: 24 base pairs

(B) TYPE: nucleotide

(C) STRANDEDNESS: single

(D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 106:

30 AGGAGTAAGG AAACCCAACG GACA

24

(2) INFORMATION FOR SEQ ID NO: 107

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 25 base pairs

35 (B) TYPE: nucleotide

(C) STRANDEDNESS: single

181

- (D) TOPOLOGY: linear  
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 107:  
TGTATATAAT GGTCTGGCTA TTGGG 25
- 5 (2) INFORMATION FOR SEQ ID NO: 108  
(i) SEQUENCE CHARACTERISTICS:  
(A) LENGTH: 26 base pairs  
(B) TYPE: nucleotide  
(C) STRANDEDNESS: single  
10 (D) TOPOLOGY: linear  
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 108:  
TTCGGCAGAA ACCTGTTATG CCAAGG 26
- (2) INFORMATION FOR SEQ ID NO: 109  
15 (i) SEQUENCE CHARACTERISTICS:  
(A) LENGTH: 26 base pairs  
(B) TYPE: nucleotide  
(C) STRANDEDNESS: single  
(D) TOPOLOGY: linear  
20 (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 109:  
GGCTCTGCTC ACAGGAGATT AGATAC 26
- (2) INFORMATION FOR SEQ ID NO: 110  
(i) SEQUENCE CHARACTERISTICS:  
25 (A) LENGTH: 26 base pairs  
(B) TYPE: nucleotide  
(C) STRANDEDNESS: single  
(D) TOPOLOGY: linear  
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 110:  
30 AAAGGCACCA GGGCCCTCAG TGAGGA 26
- (2) INFORMATION FOR SEQ ID NO: 111  
(i) SEQUENCE CHARACTERISTICS:  
35 (A) LENGTH: base pairs  
(B) TYPE: nucleotide  
(C) STRANDEDNESS: single

182

## (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 111:

GGTTTAAGAG TTGCACAAGT GCGCAGTC

28

5 (2) INFORMATION FOR SEQ ID NO: 112:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 310 base pairs

(B) TYPE: nucleotide

(C) STRANDEDNESS: single

10 (D) TOPOLOGY: linear

(ii) TYPE DE MOLECULE: ADNC

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 112:

GCTTATAGAA GGACCCCTAG TATGGGGTAA TCCCCTCTGG GAAACCAAGC CCCAGTACTC	60
AGCAGGAAAA ATAGAACATTAGG AAACCTCACA AGGACATACT TTCCCTCCCC CTAGATGGCT	120
15 AGCCACTGAG GAAGGAAAAA TACTTCACC TGCAGCTAAC CAACAGAAAT TACTTAAAAC	180
CCTTCACCAA ACCTTCCACT TAGGCATTGA TAGCACCCAT CAGATGGCCA AATTATTATT	240
TACTGGACCA GGCCTTTCA AAACATATCAA GAAGATAGTC AGGGGCTGTG AAGTGTGCCA	300
AAGAAATAAT	310

20 (2) INFORMATION FOR SEQ ID NO: 113:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 103 amino acids

(B) TYPE: amino acid

(C) STRANDEDNESS: single

25 (D) TOPOLOGY: linear

(ii) TYPE DE MOLECULE: peptide

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 113:

Leu Ile Glu Gly Pro Leu Val Trp Gly Asn Pro Leu Trp Glu Thr Lys			
1	5	10	15
30 Pro Gln Tyr Ser Ala Gly Lys Ile Glu Xaa Glu Thr Ser Gln Gly His			
20	25	30	
Thr Phe Leu Pro Ser Arg Trp Leu Ala Thr Glu Glu Gly Lys Ile Leu			
35	40	45	
Ser Pro Ala Ala Asn Gln Gln Lys Leu Leu Lys Thr Leu His Gln Thr			
35 50 55 60			
Phe His Leu Gly Ile Asp Ser Thr His Gln Met Ala Lys Leu Leu Phe			

183

65	70	75	80
Thr Gly Pro Gly Leu Phe Lys Thr Ile Lys Lys Ile Val Arg Gly Cys			
85	90	95	
Glu Val Cys Gln Arg Asn Asn			
5	100		

## (2) INFORMATION FOR SEQ ID NO: 114:

- (i) SEQUENCE CHARACTERISTICS:
- (A) LENGTH: 635 base pairs
  - 10 (B) TYPE: nucleotide
  - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear

(ii) TYPE DE MOLECULE: ADNC

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 114:

15 CCCTGTATCT TTAACCTCCT TGTAAAGTTT GTCTCTTCCA GAATCAAAAC TGTAAAAC	60
CAAATTGTTTC TTCAAATGGA GCACCAGATG GAGTCCATGA CTAAGATCCA CCGTGGACCC	120
CTGGACCGGC CTGCTAGCCC ATGCTCCGAT GTTAATGACA TTGAAGGCAC CCCTCCCGAG	180
GAAATCTCAA CTGCACAACC CCTACTATGC CCCAATTCA CGGGAAAGCAG TTAGAGCGGT	240
CATCAGCCAA CCTCCCCAAC AGCACTTGGG TTTTCCTGTT GAGAGGGGGG ACTGAGAGAC	300
20 AGGACTAGCT GGATTCCTA GGCCAACGAA GAATCCCTAA GCCTAGCTGG GAAGGTGACT	360
GCATCCACCT CTAAACATGG GGCTTGCAAC TTAGCTCACA CCCGACCAAT CAGAGAGCTC	420
ACTAAAATGC TAATTAGGCA AAAATAGGAG CTAAAGAAAT AGCCAATCAT CTATTGCCTG	480
AGAGCACAGC GGGAGGGACA AGGATCGGG A TATAACCCA GGCATTGAG CCGGCAACGG	540
CAACCCCTT TGGGTCCCTT CCCTTTGTAT GGGCGCTCTG TTTTCACTCT ATTTCACTCT	600
25 ATTAAATCTT GCAACTGAAA AAAAAAAA AAAAA	635

## (2) INFORMATION FOR SEQ ID NO: 115:

- (i) SEQUENCE CHARACTERISTICS:
- (A) LENGTH: 77 amino acids
  - 30 (B) TYPE: amino acid
  - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear

(ii) TYPE DE MOLECULE: peptide

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 115:

35 Pro Cys Ile Phe Asn Leu Leu Val Lys Phe Val Ser Ser Arg Ile Lys	
--	--

1

5

10

15

184

	Thr Val Lys Leu Gln Ile Val Leu Gln Met Glu His Gln Met Glu Ser		
	20	25	30
	Met Thr Lys Ile His Arg Gly Pro Leu Asp Arg Pro Ala Ser Pro Cys		
	35	40	45
5	Ser Asp Val Asn Asp Ile Glu Gly Thr Pro Pro Glu Glu Ile Ser Thr		
	50	55	60
	Ala Gln Pro Leu Leu Cys Pro Asn Ser Ala Gly Ser Ser		
	65	70	75

## 10 (2) INFORMATION FOR SEQ ID NO: 116:

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 32 base pairs
- (B) TYPE: nucleotide
- (C) STRANDEDNESS: single

15 (D) TOPOLOGY: linear

## (ii) TYPE DE MOLECULE: ADNc

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 116:

TGGGGTTCCA TTTGTAAGAC CATCTGTAGC TT

32

## 20 (2) INFORMATION FOR SEQ ID NO: 117:

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 1481 base pairs
- (B) TYPE: nucleotide
- (C) STRANDEDNESS: single

25 (D) TOPOLOGY: linear

## (ii) TYPE DE MOLECULE: ADNc

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 117:

	ATGGCCCTCC CTTATCATAC TTTTCTCTTT ACTGTTCTCT TACCCCCCTTT CGCTCTCACT	60
	GCACCCCTC CATGCTGCTG TACAACCAGT AGCTCCCTT ACCAAGAGTT TCTATGAAGA	120
30	ACGGGGCTTC CTGGAAATAT TGATGCCCA TCATATAGGA GTTTATCTAA GGGAAACTCC	180
	ACCTTCACTG CCCACACCCA TATGCCCGC AACTGCTATA ACTCTGCCAC TCTTGCATG	240
	CATGCAAATA CTCATTATTG GACAGGGAAA ATGATTAATC CTAGTTGTCC TGGAGGACTT	300
	GGAGCCACTG TCTGTTGGAC TTACTTCACC CATAACCAGTA TGTCTGATGG GGGTGGAATT	360
	CAAGGTCAGG CAAGAGAAAA ACAAGTAAAG GAAGCAATCT CCCAACTGAC CCGGGGACAT	420
35	AGCACCCCTA GCCCCTACAA AGGACTAGTT CTCTCAAAAC TACATGAAAC CCTCCGTACC	480
	CATACTCGCC TGGTGAGCCT ATTTAATACC ACCCTCACTC GGCTCCATGA GGTCTCAGCC	540

CAAAACCCTA CTAACGTGTT GATGTGCCTC CCCCTGCACT TCAGGCCATA CATTCAATC 600  
 CCTGTTCTG AACAAATGGAA CAACTTCAGC ACAGAAATAA ACACCACTTC CGTTTAGTA 660  
 GGACCTCTG TTTCCAATCT GGAAATAACC CATAACCAA ACCTCACCTG TGAAAATTT 720  
 AGCAATACTA TAGACACAAAC CAGCTCCAA TGCATCAGGT GGGTAACACC TCCCACACGA 780  
 5 ATAGTCTGCC TACCCCTCAGG AATATTTTT GTCTGTGGTA CCTCAGCCTA TCATTGTTG 840  
 AATGGCTCTT CAGAACATCTAT GTGCTTCCTC TCATTCTTAG TGCCCCCTAT GACCATCTAC 900  
 ACTGAACAAG ATTTATACAA TCATGTCGTA CCTAAGCCCC ACAACAAAAG AGTACCCATT 960  
 CTTCCCTTTG TTATCAGAGC AGGAGTGCTA GGCAGACTAG GTACTGGCAT TGGCAGTATC 1020  
 ACAACCTCTA CTCAGTTCTA CTACAAACTA TCTCAAGAAA TAAATGGTGA CATGGAACAG 1080  
 10 GTCACTGACT CCCTGGTCAC CTTGCAAGAT CAACTTAAC CCCTAGCAGC AGTAGTCCTT 1140  
 CAAAATCGAA GAGCTTTAGA CTTGCTAACCC GCCAAAAGAG GGGGAACCTG TTTATTTTA 1200  
 GGAGAAGAAC GCTGTTATTA TGTTAACAA TCCAGAATTG TCACTGAGAA AGTTAAAGAA 1260  
 ATTGGAGATC GAATACAATG TAGAGCAGAG GAGCTTCAAA ACACCGAACG CTGGGGCCTC 1320  
 CTCAGCCAAT GGATGCCCTG GGTTCTCCCC TTCTTAGGAC CTCTAGCAGC TCTAATATTG 1380  
 15 TTACTCCTCT TTGGACCCTG TATCTTTAAC CTCCTGTAA AGTTGTCTC TTCCAGAATT 1440  
 GAAGCTGTAA AGCTACAGAT GGTCTTACAA ATGGAACCCC A 1481

## (2) INFORMATION FOR SEQ ID NO: 118:

## (i) SEQUENCE CHARACTERISTICS:

- 20 (A) LENGTH: 493 amino acids  
 (B) TYPE: amino acid  
 (C) STRANDEDNESS: single  
 (D) TOPOLOGY: linear

## (ii) TYPE DE MOLECULE: peptide

## 25 (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 118:

	Met Ala Leu Pro Tyr His Thr Phe Leu Phe Thr Val Leu Leu Pro Pro		
1	5	10	15
	Phe Ala Leu Thr Ala Pro Pro Pro Cys Cys Cys Thr Thr Ser Ser Ser		
	20	25	30
30	Pro Tyr Gln Glu Phe Leu Xaa Arg Thr Arg Leu Pro Gly Asn Ile Asp		
	35	40	45
	Ala Pro Ser Tyr Arg Ser Leu Ser Lys Gly Asn Ser Thr Phe Thr Ala		
	50	55	60
	His Thr His Met Pro Arg Asn Cys Tyr Asn Ser Ala Thr Leu Cys Met		
35	65	70	75
	His Ala Asn Thr His Tyr Trp Thr Gly Lys Met Ile Asn Pro Ser Cys		80

186

	85	90	95
	Pro Gly Gly Leu Gly Ala Thr Val Cys Trp Thr Tyr Phe Thr His Thr		
	100	105	110
5	Ser Met Ser Asp Gly Gly Ile Gln Gly Gln Ala Arg Glu Lys Gln		
	115	120	125
	Val Lys Glu Ala Ile Ser Gln Leu Thr Arg Gly His Ser Thr Pro Ser		
	130	135	140
	Pro Tyr Lys Gly Leu Val Leu Ser Lys Leu His Glu Thr Leu Arg Thr		
	145	150	155
10	His Thr Arg Leu Val Ser Leu Phe Asn Thr Thr Leu Thr Arg Leu His		
	165	170	175
	Glu Val Ser Ala Gln Asn Pro Thr Asn Cys Trp Met Cys Leu Pro Leu		
	180	185	190
	His Phe Arg Pro Tyr Ile Ser Ile Pro Val Pro Glu Gln Trp Asn Asn		
15	195	200	205
	Phe Ser Thr Glu Ile Asn Thr Thr Ser Val Leu Val Gly Pro Leu Val		
	210	215	220
	Ser Asn Leu Glu Ile Thr His Thr Ser Asn Leu Thr Cys Val Lys Phe		
	225	230	235
20	Ser Asn Thr Ile Asp Thr Thr Ser Ser Gln Cys Ile Arg Trp Val Thr		
	245	250	255
	Pro Pro Thr Arg Ile Val Cys Leu Pro Ser Gly Ile Phe Phe Val Cys		
	260	265	270
	Gly Thr Ser Ala Tyr His Cys Leu Asn Gly Ser Ser Glu Ser Met Cys		
25	275	280	285
	Phe Leu Ser Phe Leu Val Pro Pro Met Thr Ile Tyr Thr Glu Gln Asp		
	290	295	300
	Leu Tyr Asn His Val Val Pro Lys Pro His Asn Lys Arg Val Pro Ile		
	305	310	315
30	320	325	335
	Leu Pro Phe Val Ile Arg Ala Gly Val Leu Gly Arg Leu Gly Thr Gly		
	340	345	350
	Ile Gly Ser Ile Thr Thr Ser Thr Gln Phe Tyr Tyr Lys Leu Ser Gln		
	355	360	365
	Glu Ile Asn Gly Asp Met Glu Gln Val Thr Asp Ser Leu Val Thr Leu		
35	Gln Asp Gln Leu Asn Ser Leu Ala Ala Val Val Leu Gln Asn Arg Arg		

187

	370	375	380
	Ala Leu Asp Leu Leu Thr Ala Lys Arg Gly Gly Thr Cys Leu Phe Leu		
385	390	395	400
	Gly Glu Glu Arg Cys Tyr Tyr Val Asn Gln Ser Arg Ile Val Thr Glu		
5	405	410	415
	Lys Val Lys Glu Ile Arg Asp Arg Ile Gln Cys Arg Ala Glu Glu Leu		
	420	425	430
	Gln Asn Thr Glu Arg Trp Gly Leu Leu Ser Gln Trp Met Pro Trp Val		
	435	440	445
10	Leu Pro Phe Leu Gly Pro Leu Ala Ala Leu Ile Leu Leu Leu Leu Phe		
	450	455	460
	Gly Pro Cys Ile Phe Asn Leu Leu Val Lys Phe Val Ser Ser Arg Ile		
	465	470	475
	Glu Ala Val Lys Leu Gln Met Val Leu Gln Met Glu Pro		
15	485	490	

## (2) INFORMATION FOR SEQ ID NO: 119:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 32 base pairs
- 20 (B) TYPE: nucleotide
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

(ii) TYPE DE MOLECULE: ADNc

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 119:

25 TCAAAATCGA AGAGCTTTAG ACTTGCTAAC CG 32

## (2) INFORMATION FOR SEQ ID NO: 120:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 1329 base pairs
- 30 (B) TYPE: nucleotide
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

(ii) TYPE DE MOLECULE: ADNc

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 120:

35 TCAAAATCGA AGAGCTTTAG ACTTGCTAAC CGCCAAAAGA GGGGAAACCT GTTTATTTT 60  
AGGGGAAGAA TGCTGTTAGT ATGTTAAC ATCTGGAATC ATTACTGAGA AAGTTAAAGA 120

AATTTGAGAT CGAATATAAT	GTAGAGCAGA GGACCTCAA AACACTGCAC	CCTGGGCCT 180
CCTCAGCAA TGGATGCCCT	GGACTCTCCC CTTCTTAGGA CCTCTAGCAG	CTATAATATT 240
TTTACTCCTC TTTGGACCCCT	GTATCTCAA CTTCCTGTGTT AAGTTTGCT CTTCCAGAA 300	
TGAAGCTGTA AAGCTACAAA	TAGTTCTTCA AATGGAACCC CAGATGCAGT CCATGACTAA 360	
5 AATCTACCGT GGACCCCTGG	ACCGGCCTGC TAGACTATGC TCTGATGTTA ATGACATTGA 420	
AGTCACCCCT CCCGAGGAAA	TCTCAACTGC ACAACCCCTA CTACACTCCA ATTCACTAGG 480	
AAGCAGTTAG AGCAGTTGTC	AGCCAACCTC CCCAACAGTA CTTGGGTTTT CCTGTTGAGA 540	
GGGTGGACTG AGAGACAGGA	CTAGCTGGAT TTCCCTAGGCT GACTAAGAAT CCCNAAGCCT 600	
ANCTGGGAAG GTGACCGCAT	CCATCTTAA ACATGGGCT TGCAACTTAG CTCACACCCG 660	
10 ACCAACATCAGA GAGCTCACTA AAATGCTAAT	CAGGAAAAA CAGGAGGTA AGCAATAGCC 720	
AATCATCTAT TGCCCTGAGAG	CACAGCGGGA AGGACAAGGA TTGGGATATA AACTCAGGCA 780	
TTCAAGCCAG CAACAGCAAC	CCCCTTGGG TCCCCCTCCA TTGTATGGG GCTCTGTTT 840	
CACTCTATTT CACTCTATTA	AATCATGCAA CTGCACTCTT CTGGTCCGTG TTTTTATGG 900	
CTCAAGCTGA GCTTTGTTC	GCCATCCACC ACTGCTGTT GCCACCGTCA CAGACCCGCT 960	
15 GCTGACTTCC ATCCCTTGG ATCCAGCAGA	GTGTCCACTG TGCTCCTGAT CCAGCGAGGT 1020	
ACCCATTGCC ACTCCCGATC	AGGCTAAAGG CTTGCCATTG TTCCCTGCATG GCTAAGTGCC 1080	
TGGGTTGTC CTAATAGAAC	TGAACACTGG TCACCTGGTT CCATGGTTCT CTTCCATGAC 1140	
CCACGGCTTC TAATAGAGCT	ATAACACTCA CCGCATGGCC CAAGATTCCA TTCCCTGGTA 1200	
TCTGTGAGGC CAAGAACCCC	AGGTCAAGAGA ANGTGAGGCT TGCCACCATT TGGGAAGTGG 1260	
20 CCCACTGCCA TTTTGGTAGC	GGCCCACAC CATCTGGGA GCTGTGGGAG CAAGGATCCC 1320	
CCAGTAACA		1329

## (2) INFORMATION FOR SEQ ID NO: 121:

## (i) SEQUENCE CHARACTERISTICS:

- 25 (A) LENGTH: 162 amino acids  
 (B) TYPE: amino acid  
 (C) STRANDEDNESS: single  
 (D) TOPOLOGY: linear

## (ii) TYPE DE MOLECULE: peptide

## 30 (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 121:

Gln Asn Arg Arg Ala Leu Asp Leu Leu Thr Ala Lys Arg Gly	Gly Thr		
1	5	10	15
Cys Leu Phe Leu Gly Glu Glu Cys Cys Xaa Tyr Val Asn Gln Ser Gly			
20	25	30	
35 Ile Ile Thr Glu Lys Val Lys Glu Ile Xaa Asp Arg Ile Xaa Cys Arg			

189

Ala Glu Asp Leu Gln Asn Thr Ala Pro Trp Gly Leu Leu Ser Gln Trp  
50 55 60  
Met Pro Trp Thr Leu Pro Phe Leu Gly Pro Leu Ala Ala Ile Ile Phe  
65 70 75 80  
5 Leu Leu Leu Phe Gly Pro Cys Ile Phe Asn Phe Leu Val Lys Phe Val  
85 90 95  
Ser Ser Arg Ile Glu Ala Val Lys Leu Gln Ile Val Leu Gln Met Glu  
100 105 110  
Pro Gln Met Gln Ser Met Thr Lys Ile Tyr Arg Gly Pro Leu Asp Arg  
10 115 120 125  
Pro Ala Arg Leu Cys Ser Asp Val Asn Asp Ile Glu Val Thr Pro Pro  
130 135 140  
Glu Glu Ile Ser Thr Ala Gln Pro Leu Leu His Ser Asn Ser Val Gly  
145 150 155 160  
15 Ser Ser

## (2) INFORMATION FOR SEQ ID NO: 122:

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 21 base pairs  
20 (B) TYPE: nucleotide  
(C) STRANDEDNESS: single  
(D) TOPOLOGY: linear

## (ii) TYPE DE MOLECULE: ADNc

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 122:

25 GGCATTGATA GCACCCATCA G 21

## (2) INFORMATION FOR SEQ ID NO: 123:

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 21 base pairs  
30 (B) TYPE: nucleotide  
(C) STRANDEDNESS: single  
(D) TOPOLOGY: linear

## (ii) TYPE DE MOLECULE: ADNc

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 123:

35 CATGTCACCA GGGTGGAATA G 21

190

## (2) INFORMATION FOR SEQ ID NO: 124:

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 758 base pairs
- (B) TYPE: nucleotide
- 5 (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

## (ii) TYPE DE MOLECULE: ADNc

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 124:

GGCATTGATA	GCACCCATCA	GATGCCAAA	TCATTATTTA	CTGGACCAGG	CCTTTCAA	60
10 ACTATCAAGC	AGATAGGGCC	CGTGAAGCAT	GCCAAAGAAA	TAATCCCCTG	CCTTATGCC	120
ATGTTCTTC	AGGAGAACAA	AGAACAGGCC	ATTACCCAGG	GGAAGACTGG	CAACTAGATT	180
TTACCCACAT	GGCCAAATGT	CAGGGATTTC	AGCATCTACT	AGTCTGGCA	GATACTTCA	240
CTGGTTGGGT	GGAGTCTTCT	CCTTGTAGGA	CAGAAAAGAC	CCAAGAGGTA	ATAAAGGCAC	300
TAATGAAATA	ATTCCCAGAT	TTGGACTTCC	CCCAGGATTA	CAGGGTGACA	ATGGCCCCGC	360
15 TTTCAAGGCT	GCAGTAACCC	AGGGAGTATC	CCAGGTGTTA	GGCATACAAT	ATCACTTACA	420
CTGTGCCTGG	AGGCCACAAT	CCTCCAGAAA	AGTCAGAAA	ATGAATGAAA	CACTCAAAGA	480
TCTAAAAAAAG	CTAACCAAG	AAACCCACAT	TGCATGACCT	GTTCTGTTGC	CTATAACCTT	540
ACTAAGAATC	CATAACTATC	CCCCAAAAAG	CAGGACTTAG	CCCATACGAG	ATGCTATATG	600
GATGGCCTTT	CCTAACCAAT	GACCTTGTC	TTGACTGAGA	AATGGCCAAC	TTAGTTGCAG	660
20 ACATCACCTC	CTTAGCCAAA	TATCAACAAG	TTCTTAAAC	ATCACAGGGA	ACCTGTCCCC	720
GAGAGGAGGG	AAAGGAACTA	TTCCACCCCTG	GTGACATG			758

## (2) INFORMATION FOR SEQ ID NO: 126:

## 25 (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 25 base pairs
- (B) TYPE: nucleotide
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

## 30 (ii) TYPE DE MOLECULE: ADNc

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 126:

CGGACATCCA AAGTGATGGG AAACG

25

## (2) INFORMATION FOR SEQ ID NO: 127:

## 35 (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 26 base pairs

191

(B) TYPE: nucleotide  
(C) STRANDEDNESS: single  
(D) TOPOLOGY: linear  
(ii) TYPE DE MOLECULE: ADNc  
5 (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 127:  
GGACAGGAAA GTAAGACTGA GAAGGC

26

(2) INFORMATION FOR SEQ ID NO: 128:  
(i) SEQUENCE CHARACTERISTICS:  
10 (A) LENGTH: 26 base pairs  
(B) TYPE: nucleotide  
(C) STRANDEDNESS: single  
(D) TOPOLOGY: linear  
(ii) TYPE DE MOLECULE: ADNc  
15 (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 128:  
CCTAGAACGT ATTCTGGAGA ATTGGG

26

(2) INFORMATION FOR SEQ ID NO: 129:  
(i) SEQUENCE CHARACTERISTICS:  
20 (A) LENGTH: 26 base pairs  
(B) TYPE: nucleotide  
(C) STRANDEDNESS: single  
(D) TOPOLOGY: linear  
(ii) TYPE DE MOLECULE: ADNc  
25 (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 129:  
TGGCTCTCAA TGGTCAAACA TACCCG

26

(2) INFORMATION FOR SEQ ID NO: 130:  
(i) SEQUENCE CHARACTERISTICS:  
30 (A) LENGTH: 1511 base pairs  
(B) TYPE: nucleotide  
(C) STRANDEDNESS: single  
(D) TOPOLOGY: linear  
(ii) TYPE DE MOLECULE: ADNc  
35 (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 130:  
CCTAGAACGT ATTCTGGAGA ATTGGGACCA ATGTGACACT CAGACGCTAA GAAAGAAACC

60

192

	ATTTATATTC TTCTGCAGTA CCGCCTGGCC ACAATATCCT CTTCAAGGGA GAGAAACCTG	120
	GCTTCCTGAG GGAAGTATAA ATTATAACAT CATCTTACAG CTAGACCTCT TCTGTAGAAA	180
	GGAGGGCAAA TGGAGTGAAG TGCCATATGT GCAAACTTTC TTTTCATTAA GAGACAACTC	240
	ACAATTATGT AAAAAGTGTG GTTTATGCC CACAGGAAGC CCTCAGAGTC CACCTCCCTA	300
5	CCCCAGCGTC CCCTCCCGA CTCCTTCCTC AACTAATAAG GACCCCCCTT TAACCCAAAC	360
	GGTCCAAAAG GAGATAGACA AAGGGTAAA CAATGAACCA AAGAGTGCCA ATATTCCCCG	420
	ATTATGCCCTC CTCCAAGCAG TGAGAGGAGG AGAATTCCGC CCAGCCAGAG TGCCGTAC	480
	TTTTTCTCTC TCAGACTTAA AGCAAATTAA AATAGACCTA GGTAAATTCT CAGATAACCC	540
	TGACGGCTAT ATTGATGTT TACAAGGTT AGGACAATCC TTTGATCTGA CATGGAGAGA	600
10	TATAATGTTA CTACTAAATC AGACACTAAC CCCAAATGAG AGAAGTGCCG CTGTAAC	660
	AGCCCGAGAG TTTGGCGATC TTTGGTATCT CAGTCAGGCC AACAAATAGGA TGACAACAGA	720
	GGAAAGAACAA ACTCCCACAG GCCAGCAGGC AGTCCCAGT GTAGACCCCTC ATTGGGACAC	780
	AGAATCAGAA CATGGAGATT GGTGCCACAA ACATTTGCTA ACTTGCCTGC TAGAAGGACT	840
	GAGGAAAAACT AGGAAGAACG CTATGAATT CTCAATGATG TCCACTATAA CACAGGGAAA	900
15	GGAAGAAAAT CTTACTGCTT TTCTGGACAG ACTAAGGGAG GCATTGAGGA AGCATAC	960
	CCTGTCACCT GACTCTATTG AAGGCCACT AATCTAAAG GATAAGTTA TCACTCAGTC	1020
	AGCTGCAGAC ATTAGAAAAA ACTTCAAAAG TCTGCCTTAG GCCCGGAGCA GAACTTAGAA	1080
	ACCCATTTA ACTTGGCATC CTCAGTTTT TATAATAGAG ATCAGGAGGA GCAGGGAAA	1140
	CGGGACAAAC GGGATAAAAAA AAAAAGGGGG GGTCCACTAC TTTAGTCATG GCCCTCAGGC	1200
20	AAGCAGACTT TGGAGGCTCT GCaaaAGGGA AAAGCTGGC AAATCAAATG CCTAATAGGG	1260
	CTGGCTTCCA GTGCGGTCTA CAAGGACACT TTAAAAAAGA TTATCCAAGT AGAAATAAGC	1320
	CGCCCCCTTG TCCATGCCCT TTACGTCAAG GGAATCACTG GAAGGCCAC TGCCCCAGGG	1380
	GATGAAGATA CTCTGAGTCA GAAGCCATTA ACCAGATGAT CCAGCAGCAG GACTGAGGGT	1440
	GCCCCGGGCG AGCGCCAGCC CATGCCATCA CCCTCACAGA GCCCCGGGTA TGTTGACCA	1500
25	TTGAGAGCCA A	1511

## (2) INFORMATION FOR SEQ ID NO: 131:

## (i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 352 amino acids

30 (B) TYPE: amino acid

(C) STRANDEDNESS: single

(D) TOPOLOGY: linear

(ii) TYPE DE MOLECULE: peptide

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 131:

35 Leu Glu Arg Ile Leu Glu Asn Trp Asp Gln Cys Asp Thr Gln Thr Leu

193

Arg Lys Lys Arg Phe Ile Phe Phe Cys Ser Thr Ala Trp Pro Gln Tyr  
20 25 30  
Pro Leu Gln Gly Arg Glu Thr Trp Leu Pro Glu Gly Ser Ile Asn Tyr  
35 40 45  
5 Asn Ile Ile Leu Gln Leu Asp Leu Phe Cys Arg Lys Glu Gly Lys Trp  
50 55 60  
Ser Glu Val Pro Tyr Val Gln Thr Phe Phe Ser Leu Arg Asp Asn Ser  
65 70 75 80  
Gln Leu Cys Lys Lys Cys Gly Leu Cys Pro Thr Gly Ser Pro Gln Ser  
10 85 90 95  
Pro Pro Pro Tyr Pro Ser Val Pro Ser Pro Thr Pro Ser Ser Thr Asn  
100 105 110  
Lys Asp Pro Pro Leu Thr Gln Thr Val Gln Lys Glu Ile Asp Lys Gly  
115 120 125  
15 Val Asn Asn Glu Pro Lys Ser Ala Asn Ile Pro Arg Leu Cys Pro Leu  
130 135 140  
Gln Ala Val Arg Gly Gly Glu Phe Gly Pro Ala Arg Val Pro Val Pro  
145 150 155 160  
Phe Ser Leu Ser Asp Leu Lys Gln Ile Lys Ile Asp Leu Gly Lys Phe  
20 165 170 175  
Ser Asp Asn Pro Asp Gly Tyr Ile Asp Val Leu Gln Gly Leu Gly Gln  
180 185 190  
Ser Phe Asp Leu Thr Trp Arg Asp Ile Met Leu Leu Leu Asn Gln Thr  
195 200 205  
25 Leu Thr Pro Asn Glu Arg Ser Ala Ala Val Thr Ala Ala Arg Glu Phe  
210 215 220  
Gly Asp Leu Trp Tyr Leu Ser Gln Ala Asn Asn Arg Met Thr Thr Glu  
225 230 235 240  
Glu Arg Thr Thr Pro Thr Gly Gln Gln Ala Val Pro Ser Val Asp Pro  
30 245 250 255  
His Trp Asp Thr Glu Ser Glu His Gly Asp Trp Cys His Lys His Leu  
260 265 270  
Leu Thr Cys Val Leu Glu Gly Leu Arg Lys Thr Arg Lys Lys Pro Met  
275 280 285  
35 Asn Tyr Ser Met Met Ser Thr Ile Thr Gln Gly Lys Glu Glu Asn Leu  
290 295 300

194

Thr Ala Phe Leu Asp Arg Leu Arg Glu Ala Leu Arg Lys His Thr Ser  
305 310 315 320  
Leu Ser Pro Asp Ser Ile Glu Gly Gln Leu Ile Leu Lys Asp Lys Phe  
325 330 335  
5 Ile Thr Gln Ser Ala Ala Asp Ile Arg Lys Asn Phe Lys Ser Leu Pro  
340 345 350

## (2) INFORMATION FOR SEQ ID NO: 132:

## (i) SEQUENCE CHARACTERISTICS:

- 10 (A) LENGTH: 30 base pairs  
(B) TYPE: nucleotide  
(C) STRANDEDNESS: single  
(D) TOPOLOGY: linear

(ii) TYPE DE MOLECULE: ADNc

15 (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 132:

TGCTGGAATT CGGGATCCTA GAACGTATTC

30

## (2) INFORMATION FOR SEQ ID NO: 133:

## (i) SEQUENCE CHARACTERISTICS:

- 20 (A) LENGTH: 30 base pairs  
(B) TYPE: nucleotide  
(C) STRANDEDNESS: single  
(D) TOPOLOGY: linear

(ii) TYPE DE MOLECULE: ADNc

25 (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 133:

AGTTCTGCTC CGAAGCTTAG GCAGACTTTT

30

## (2) INFORMATION FOR SEQ ID NO: 135:

## (i) SEQUENCE CHARACTERISTICS:

- 30 (A) LENGTH: 398 amino acids  
(B) TYPE: amino acid  
(C) STRANDEDNESS: single  
(D) TOPOLOGY: linear

(ii) TYPE DE MOLECULE: peptide

35 (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 135:

Met Gly Ser Ser His His His His His His Ser Ser Gly Leu Val Pro

195

1	5	10	15
Arg Gly Ser His Met Ala Ser Met Thr Gly Gly Gln Gln Met Gly Arg			
20	25	30	
Ile Leu Glu Arg Ile Leu Glu Asn Trp Asp Gln Cys Asp Thr Gln Thr			
5	35	40	45
Leu Arg Lys Lys Arg Phe Ile Phe Phe Cys Ser Thr Ala Trp Pro Gln			
50	55	60	
Tyr Pro Leu Gln Gly Arg Glu Thr Trp Leu Pro Glu Gly Ser Ile Asn			
65	70	75	80
10	Tyr Asn Ile Ile Leu Gln Leu Asp Leu Phe Cys Arg Lys Glu Gly Lys		
	85	90	95
Trp Ser Glu Val Pro Tyr Val Gln Thr Phe Phe Ser Leu Arg Asp Asn			
	100	105	110
Ser Gln Leu Cys Lys Cys Gly Leu Cys Pro Thr Gly Ser Pro Gln			
15	115	120	125
Ser Pro Pro Pro Tyr Pro Ser Val Pro Ser Pro Thr Pro Ser Ser Thr			
	130	135	140
Asn Lys Asp Pro Pro Leu Thr Gln Thr Val Gln Lys Glu Ile Asp Lys			
145	150	155	160
20	Gly Val Asn Asn Glu Pro Lys Ser Ala Asn Ile Pro Arg Leu Cys Pro		
	165	170	175
Leu Gln Ala Val Arg Gly Gly Glu Phe Gly Pro Ala Arg Val Pro Val			
	180	185	190
Pro Phe Ser Leu Ser Asp Leu Lys Gln Ile Lys Ile Asp Leu Gly Lys			
25	195	200	205
Phe Ser Asp Asn Pro Asp Gly Tyr Ile Asp Val Leu Gln Gly Leu Gly			
	210	215	220
Gln Ser Phe Asp Leu Thr Trp Arg Asp Ile Met Leu Leu Leu Asn Gln			
225	230	235	240
30	Thr Leu Thr Pro Asn Glu Arg Ser Ala Ala Val Thr Ala Ala Arg Glu		
	245	250	255
Phe Gly Asp Leu Trp Tyr Leu Ser Gln Ala Asn Asn Arg Met Thr Thr			
	260	265	270
Glu Glu Arg Thr Thr Pro Thr Gly Gln Gln Ala Val Pro Ser Val Asp			
35	275	280	285
Pro His Trp Asp Thr Glu Ser Glu His Gly Asp Trp Cys His Lys His			

196

290	295	300
Leu Leu Thr Cys Val Leu Glu Gly Leu Arg Lys Thr Arg Lys Lys Pro		
305	310	315
Met Asn Tyr Ser Met Met Ser Thr Ile Thr Gln Gly Lys Glu Glu Asn		
5	325	330
Leu Thr Ala Phe Leu Asp Arg Leu Arg Glu Ala Leu Arg Lys His Thr		
340	345	350
Ser Leu Ser Pro Asp Ser Ile Glu Gly Gln Leu Ile Leu Lys Asp Lys		
355	360	365
10 Phe Ile Thr Gln Ser Ala Ala Asp Ile Arg Lys Asn Phe Lys Ser Leu		
370	375	380
Pro Lys Leu Ala Ala Ala Leu Glu His His His His His His His		
385	390	395

## 15 (2) INFORMATION FOR SEQ ID NO: 137:

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 378 amino acids
- (B) TYPE: amino acid
- (C) STRANDEDNESS: single
- 20 (D) TOPOLOGY: linear

## (ii) TYPE DE MOLECULE: peptide

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 137:

Met Ala Ser Met Thr Gly Gly Gln Gln Met Gly Arg Ile Leu Glu Arg		
1	5	10
25 Ile Leu Glu Asn Trp Asp Gln Cys Asp Thr Gln Thr Leu Arg Lys Lys		
20	25	30
Arg Phe Ile Phe Phe Cys Ser Thr Ala Trp Pro Gln Tyr Pro Leu Gln		
35	40	45
Gly Arg Glu Thr Trp Leu Pro Glu Gly Ser Ile Asn Tyr Asn Ile Ile		
30 50	55	60
Leu Gln Leu Asp Leu Phe Cys Arg Lys Glu Gly Lys Trp Ser Glu Val		
65	70	75
Pro Tyr Val Gln Thr Phe Phe Ser Leu Arg Asp Asn Ser Gln Leu Cys		
85	90	95
35 Lys Lys Cys Gly Leu Cys Pro Thr Gly Ser Pro Gln Ser Pro Pro Pro		
100	105	110

197

Tyr Pro Ser Val Pro Ser Pro Thr Pro Ser Ser Thr Asn Lys Asp Pro  
115 120 125  
Pro Leu Thr Gln Thr Val Gln Lys Glu Ile Asp Lys Gly Val Asn Asn  
130 135 140  
5 Glu Pro Lys Ser Ala Asn Ile Pro Arg Leu Cys Pro Leu Gln Ala Val  
145 150 155 160  
Arg Gly Gly Glu Phe Gly Pro Ala Arg Val Pro Val Pro Phe Ser Leu  
165 170 175  
Ser Asp Leu Lys Gln Ile Lys Ile Asp Leu Gly Lys Phe Ser Asp Asn  
10 180 185 190  
Pro Asp Gly Tyr Ile Asp Val Leu Gln Gly Leu Gly Gln Ser Phe Asp  
195 200 205  
Leu Thr Trp Arg Asp Ile Met Leu Leu Leu Asn Gln Thr Leu Thr Pro  
210 215 220  
15 Asn Glu Arg Ser Ala Ala Val Thr Ala Ala Arg Glu Phe Gly Asp Leu  
225 230 235 240  
Trp Tyr Leu Ser Gln Ala Asn Asn Arg Met Thr Thr Glu Glu Arg Thr  
245 250 255  
Thr Pro Thr Gly Gln Gln Ala Val Pro Ser Val Asp Pro His Trp Asp  
20 260 265 270  
Thr Glu Ser Glu His Gly Asp Trp Cys His Lys His Leu Leu Thr Cys  
275 280 285  
Val Leu Glu Gly Leu Arg Lys Thr Arg Lys Lys Pro Met Asn Tyr Ser  
290 295 300  
25 Met Met Ser Thr Ile Thr Gln Gly Lys Glu Glu Asn Leu Thr Ala Phe  
305 310 315 320  
Leu Asp Arg Leu Arg Glu Ala Leu Arg Lys His Thr Ser Leu Ser Pro  
325 330 335  
Asp Ser Ile Glu Gly Gln Leu Ile Leu Lys Asp Lys Phe Ile Thr Gln  
30 340 345 350  
Ser Ala Ala Asp Ile Arg Lys Asn Phe Lys Ser Leu Pro Lys Leu Ala  
355 360 365  
Ala Ala Leu Glu His His His His His His  
370 375

35

(2) INFORMATION FOR SEQ ID NO: 138:

198

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 25 base pairs
- (B) TYPE: nucleotide
- (C) STRANDEDNESS: single

5 (D) TOPOLOGY: linear

## (ii) TYPE DE MOLECULE: ADNc

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 138:

CTTGGAGGGT GCATAACCA GGAAT

25

10 (2) INFORMATION FOR SEQ ID NO: 139:

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 20 base pairs
- (B) TYPE: nucleotide
- (C) STRANDEDNESS: single

15 (D) TOPOLOGY: linear

## (ii) TYPE DE MOLECULE: ADNc

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 139:

TGTCCGCTGT GCTCCTGATC

20

20 (2) INFORMATION FOR SEQ ID NO: 140:

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 25 base pairs
- (B) TYPE: nucleotide
- (C) STRANDEDNESS: single

25 (D) TOPOLOGY: linear

## (ii) TYPE DE MOLECULE: ADNc

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 140:

CTATGTCCTT TTGGACTGTT TGGGT

25

30 (2) INFORMATION FOR SEQ ID NO: 141:

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 764 base pairs
- (B) TYPE: nucleotide
- (C) STRANDEDNESS: single

35 (D) TOPOLOGY: linear

## (ii) TYPE DE MOLECULE: ADNc

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 141:

TGTCCGCTGT	GCTCCTGATC	CAGCACAGGC	GCCCATTGCC	TCTCCAATT	GGGCTAAAGG	60	
CTTGCCATTG	TTCCTGCACA	GCTAAGTGCC	TGGGTTCATC	CTAATCGAGC	TGAACACTAG	120	
TCACTGGGTT	CCACGGTTCT	CTTCCATGAC	CCATGGCTTC	TAATAGAGCT	ATAACACTCA	180	
5	CTGCATGGTC	CAAGATTCCA	TTCCTTGAA	TCCGTGAGAC	CAAGAACCCC	AGGTCAAGAGA	240
	ACACAAGGCT	TGCCACCATG	TTGGAAGCAG	CCCACCACCA	TTTGGAAAGC	AGCCCCCAC	300
	TATCTTGGGA	GCTCTGGGAG	CAAGGACCCC	AGGTAACAAT	TTGGTGCACCA	CGAAGGGGAC	360
	TGAATCCGCA	ACCATGAAGG	GATCTCCAAA	GCAATTGGAA	ATGTTCTCC	CAAGGCAAAA	420
	ATGCCCTAA	GATGTATTCT	GGAGAATTGG	GACCAATTG	ACCCCTCAGAC	AGTAAGAAAA	480
10	AAATGACTTA	TATTCTTCTG	CAGTACCGCC	CTGGCCACGA	TATCCTCTTC	AAGGGGGAGA	540
	AACCTGGCCT	CCTGAGGGAA	GTATAAATTA	TAACACCATC	TTACAGCTAG	ACCTGTTTG	600
	TAGAAAAGGA	GGCAAATGGA	GTGAAGTGCC	ATATTACAA	ACTTTCTTT	CATTAAAAGA	660
	CAACTCGCAA	TTATGTTAAC	AGTGTGATT	GTGTTCTAC	ACGGAAGCCC	TCAGATTCTA	720
	CTCCCCACCC	CGGCATCTC	CCCTGAATCC	CTCCCCAACT	TATT		764

15

## (2) INFORMATION FOR SEQ ID NO: 142:

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 800 base pairs
- (B) TYPE: nucleotide
- 20 (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

## (ii) TYPE DE MOLECULE: ADNC

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 142:

TGTCCGCTGT	GCTCCTGATC	CAGCACAGGC	GCCCATTGCC	TCTCCAATT	GGGCTAAAGG	60	
25	CTTGCCATTG	TTCCTGCACA	GCTAAGTGCC	TGGGTTCATC	CTAATCGAGC	TGAACACTAG	120
	TCACTGGGTT	CCACGGTTCT	CTTCCATGAC	CCATGGCTTC	TAATAGAGCT	ATAACACTCA	180
	CTGCATGGTC	CAAGATTCCA	TTCCTTGAA	TCCGTGAGAC	CAAGAACCCC	AGGTCAAGAGA	240
	ACACAAGGCT	TGCCACCATG	TTGGAAGCAG	CCCACCACCA	TTTGGAAAGC	GGCCCCCAC	300
	TATCTTGGGA	GCTCTGGGAG	CAAGGACCCC	CAGGTAACAA	TTGGTGACC	ACGAAGGGAC	360
30	CTGAATCCGC	AACCATGAAG	GGATCTCCAA	AGCAATTGGA	AATGTTCTC	CCAAGGCAAA	420
	AATGCCCTA	AGATGTATT	TGGAGAATTG	GGACCAATCT	GACCTCAGA	CAGTAAGAAA	480
	AAAAATGACT	TATATTCTTC	TGCAGTACCG	CCTGGCCACG	GATATCCTCT	TCAAGGGGGA	540
	GAAACCTGGC	CTCCTGAGGG	AAGTATAAAT	TATAACACCA	TCTTACAGCT	AGACCTGTT	600
	TGTAGAAAAG	GAGGCAAATG	GAGTGAAGTG	CCATATTAC	AAACTTCTT	TTCATTAAGA	660
35	GACAACTCGC	AATTATGTAA	ACAGTGTGAT	TTGTGTCCTA	CAGGAAGCCC	TCAGATCTAC	720
	CTCCCTACCC	CGGCATCTC	CTGACTCCTT	CCCCAACTAA	TAAGGACCCA	CTTCAGCCCA	780

200

AACAGTCCAA AAGGACATAG

800

## (2) INFORMATION FOR SEQ ID NO: 169:

## (i) SEQUENCE CHARACTERISTICS:

- 5           (A) LENGTH: base pairs  
             (B) TYPE: nucleotide  
             (C) STRANDEDNESS: single  
             (D) TOPOLOGY: linear

(ii) TYPE DE MOLECULE: ADNc

10           (xi) SEQUENCE DESCRIPTION: SEQ ID NO:169:

consensus (41/68-1 + 42/68-1 + c143 68-1)

## (2) INFORMATION FOR SEQ ID NO: 170:

## (i) SEQUENCE CHARACTERISTICS:

- 15           (A) LENGTH: 438 base pairs  
             (B) TYPE: nucleotide  
             (C) STRANDEDNESS: single  
             (D) TOPOLOGY: linear

(ii) TYPE DE MOLECULE: ADNc

20           (xi) SEQUENCE DESCRIPTION: SEQ ID NO:170:

GA	CTTGAGCC	AGTCCTCATA	CCTGGACACT	CTTGTCTTC	GGTACATGGA	TGATTACTT	60
TT	AGGCCACCC	ATTCAGAAC	CTTGTGCCAT	CAAGCCACCC	AAGCACTCTT	AAATTCCTT	120
G	GCTACCTGTG	GCTACAAGGT	TTCCAAACCA	AAGGCTCAGC	TCTGCTCACA	GCAGGTTAAA	180
T	TACTTAGGGC	TAAAATTATC	CAAAGGCACC	AGAACCCCTCA	GTGAGGAACG	TATCCAGCCT	240
25	ATACTGGGTT	ATCCTCATCC	CAAAACCTA	AAGCAACTAA	CAGCGTTCCCT	TGGCATAACA	300
GG	GGTTCTGCC	AAATATGGAT	TCCCAGGTAC	ACCAAGATAG	CCAGACCATT	AAATACACGA	360
AT	ATTAAGGAAA	CTCAAAAGC	CAATACCCAT	TTAGTAAGAT	GGACACCTGA	AGCAGAAGTG	420
G	GCTTCCAGG	CCCTAAAG					438

30           (2) INFORMATION FOR SEQ ID NO: 171:

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 438 base pairs  
             (B) TYPE: nucleotide  
             (C) STRANDEDNESS: single

35           (D) TOPOLOGY: linear

(ii) TYPE DE MOLECULE: ADNc

201

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO:171:

	GACTTGAGCC AGTCCTCATA CCTGGACACT CTTGTCCTTC GGTACATGGA TGATTACTT	60
	TTAGGCCACCC ATTCAAGAAC CTTGTGCCAT CAAGCCACCC AAGCACTCTT AAATTCCTT	120
	GCTACCTGTG GCTACAAGGT TTCCAAACCA AAGGCTCAGC TCTGCTCACA GCAGGTTAAA	180
5	TACTTAGGGC TAAAATTATC CAAAGGCACC AGAACCTCTA GTGAGGAACG TATCCAGCCT	240
	ATACTGGGTT ATCCTCATCC CAAAACCCCTA AAGCAACTAA CAGCGTTCCCT TGGCATAACA	300
	GGTTTCTGCC AAATATGGAT TCCCAGGTAC AGCAAAGTAG CCAGACCATT AAATACACGA	360
	ATTAAGGAAA CTCAAAAAGC CAGTACCCAT TTAGTAAGAT GGACACCTGA AGCAGAAGTG	400
	GCTTCCAGG CCCTAAAG	438

10

## (2) INFORMATION FOR SEQ ID NO: 172:

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 438 base pairs
- (B) TYPE: nucleotide
- 15 (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

## (ii) TYPE DE MOLECULE: ADNC

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO:172:

	GACTTGAGCC AGTCYTCATA CCTGGACAYT CTTGTCCTTC GGTACATGGA TGATTACTT	60
20	TTAGGCCACCC ATTCAAGAAC CTTGTGCCAT CAAGCCACCC AAGCACTCTT AAATTCCTT	120
	GCTACCTGTG GCTACAAGGT TTCCAAACCA AAGGCTCAGC TCTGCTCACA GCAGGTTAAA	180
	TACTTAGGGC TAAAATTATC CAAAGGCACC AGAACCTCTA GTGAGGAACG TATCCAGCCT	240
	ATACTGGGTT ATCCTCATCC CAAAACCCCTA AAGCAACTAA CAGCGTTCCCT TGGCATAACA	300
	GGTTTCTGCC AAATATGGAT TCCCAGGTAC AGCAAATAG CCAGACCATT AAATACACGA	360
25	ATTAAGGAAA CTCAAAAAGC CAATACCCAT TTAGTAAGAT GGACATCTGA AGCAGAAGTG	400
	GCTTCCAGG CCCTAAAG	438

## (2) INFORMATION FOR SEQ ID NO: 173:

## (i) SEQUENCE CHARACTERISTICS:

- 30 (A) LENGTH: 146 amino acids
- (B) TYPE: amino acid
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

## (ii) TYPE DE MOLECULE: peptide

## 35 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:173:

DLSQSSYLDL LVLRYMDDLL LATHSETLCH QATQALLNFL ATCGYKVSKP

50

KAQLCSQQVK YLGLKLSKGT RTLSEERIQP ILGYPHPKTL KQLTAFLGIT	100
GFCQIWIPRY SKIARPLNTR IKETQKANTH LVRWTPEAEV AFQALK	146

## (2) INFORMATION FOR SEQ ID NO: 174:

5 (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 146 amino acids
- (B) TYPE: amino acid
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

10 (ii) TYPE DE MOLECULE: peptide

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:174:

DLSQSSYLDT LVLRYMDDLL LATHSETLCH QATQALLNFL ATCGYKVSKP	50
KAQLCSQQVK YLGLKLSKGT RTLSEERIQP ILGYPHPKTL KQLTAFLGIT	100
GFCQIWIPRY SKVARPLNTR IKETQKASTH LVRWTPEAEV AFQALK	146

15

## (2) INFORMATION FOR SEQ ID NO: 175:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 146 amino acids
- (B) TYPE: amino acid
- 20 (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

(ii) TYPE DE MOLECULE: peptide

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:175:

DLSQSSYLDX LVLRYMDDLL LATHSETLCH QATQALLNFL ATCGYKVSKP	50
25 KAQLCSQQVK YLGLKLSKGT RTLSEERIQP ILGYPHPKTL KQLTAFLGIT	100
GFCQIWIPRY SKIARPLNTR IKETQKANTH LVRWTSEAEV AFQALK	146

## (2) INFORMATION FOR SEQ ID NO: 176:

(i) SEQUENCE CHARACTERISTICS:

30 (A) LENGTH: base pairs
(B) TYPE: nucleotide
(C) STRANDEDNESS: single
(D) TOPOLOGY: linear

(ii) TYPE DE MOLECULE: ADNC

35 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:176:

consensus (1/46-7+8/46-7+c15/46/7)

## (2) INFORMATION FOR SEQ ID NO: 177:

## (i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 429 base pairs

5

(B) TYPE: nucleotide

(C) STRANDEDNESS: single

(D) TOPOLOGY: linear

(ii) TYPE DE MOLECULE: ADNc

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:177:

10	GACTTGAGCC AGTCCTCATA CCTGGACATT CTTGTTCTTC AGTATGGGA TGACTTAATT	60
	ATAGCCACCC ATTCAAGAAC CTTGTGGCAT CAAGCCACCC AAGCGCTCTT AAATTCCTT	120
	GCTACCTGTG GCTCCAAACA AAAGGCTCAC CTCTGCTCAC ACCAGGTTAA ATACTTAGGG	180
	CTAAAATTAT CCAAAGTCAC CAGGGCCCTC AGAGAGGAAC GTATCCAGCG TATACTGGCT	240
	TATCCTCATC CCATAACCCCT AAAGCAACTA AGAGGGTTCC TTGGCATATC AGCCTTCTGC	300
15	CGAATATGGA TTCCCCGATA CAGTGAAATA GCCAGGCCAT TATGTACATT AATTAAGGAA	360
	ACTCAGAAAG CCAATACCCA TATAGTAAGA TGGACACCTG AAACAGAAGT GGCTTCCAG	420
	GCCCTAAAG	429

## (2) INFORMATION FOR SEQ ID NO: 178:

## (i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 429 base pairs

(B) TYPE: nucleotide

(C) STRANDEDNESS: single

(D) TOPOLOGY: linear

25

(ii) TYPE DE MOLECULE: ADNc

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:178:

	GACTTGAGCC AGTCCTCATA CCTGGACATT CTTGTTCTTC AGTATAGGGA TGATTTAATT	60
	ATAGCCACCC ATTCAAGAAC CTTGTGGCAT CAAGCCACCC AAGTGCTCTT AAATTCCTC	120
	GCTACCTGTG GCTCCAAACA AAGGGCTCAC CTCTGCTCAC AGCAGGTTAA ATACTTAGGG	180
30	CTAAAATTAT CCAAAGTCGC CAGGGCCCTC AGAGAGGAAC GTATCCAGCG TATACTGGAT	240
	TATCCTCATC CCAAAACCAT AAAGCAACTA AGAGGGTTCC TTGGCATAAC AGCCTTCTGC	300
	CGAATATGGA TTCCCCGATA CAGTGAAATA GCCAGGCCAT TATGTACATT AGTTAAGGAA	360
	ACTCAGAAAG CCAATACCCA TATAGTAAGA TGGACACCTG AGACAGAAGT GGCTTCCAG	420
	GCCCTAAAG	429

35

## (2) INFORMATION FOR SEQ ID NO: 179:

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 429 base pairs
- (B) TYPE: nucleotide
- (C) STRANDEDNESS: single

5                   (D) TOPOLOGY: linear

## (ii) TYPE DE MOLECULE: ADNc

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO:179:

GACTTGAGCC	AGTCCTCATA	CCTGGACATT	CTTGTTCCTC	AGTATGGGGA	TGATTTAATT	60
ATAGCCACCC	ATTCAAGAAC	CTTGTGGCAC	CAAGCCACCC	AAGCGCTCTT	AAATTCCTC	120
10 GCTACCTGTG	GCTCCAAACA	AAAGGCTCAG	CTCTGCTCAC	AGCAGGTTAA	ATACTTAGGG	180
CTAAAATTAT	CCAAAGTCAC	CAGGGCCCTC	AGAGAGGAAC	GTATCCAGCG	TATACTGGCT	240
TATCCCCATC	CCAAACCCCT	AAAGCAACTA	AGARGGTTCC	TTGGCATAAC	AGCCTTCTGC	300
CGAATATGGA	TTCCCGAGATA	CAGCGAAATA	GCCAGGCCAT	TATGTACATT	ATCTAAGGAA	360
ACTCAGAAAG	CCAATACCCA	TATAGTAAGA	TGGACACCTG	AAACAGAAAGT	GGCTTCCAG	420
15 GCCCTAAAG						429

## (2) INFORMATION FOR SEQ ID NO: 180:

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 143 amino acids
- 20 (B) TYPE: amino acid
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

## (ii) TYPE DE MOLECULE: peptide

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO:180:

DLSQSSYLDI	LVLQYGDLLI	IATHSETLWH	QATQALLNFL	ATCGSKQKAH	50
LCSHQVKYLG	LKLSKVTRAL	REERIQRILA	YPHPITLKQL	RGFLGISAFC	100
RIWIPGYSEI	ARPLCLIKE	TQKANTHIVR	WTPETEVAFQ	ALK	143

## (2) INFORMATION FOR SEQ ID NO: 181:

30                   (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 143 amino acids
- (B) TYPE: amino acid
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

35                   (ii) TYPE DE MOLECULE: peptide

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO:181:

205

DLSQSSYLDI LVLQYRDDL IATHSETLWH QATQVLLNFL ATCGSKQRAQ	50
LCSQQVKYLG LKLSKVARAL REERIQRILD YPHPKTIKQL RGFLGITAF	100
RIWIPRYSEI ARPLCTLVKE TQKANTHIVR WTPETEVAFQ ALK	143

## 5 (2) INFORMATION FOR SEQ ID NO: 182:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 143 amino acids
  - (B) TYPE: amino acid
  - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear

(ii) TYPE DE MOLECULE: peptide

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:182:

DLSQSSYLDI LVPQYGDDLI IATHSETLWH QATQALLNFL ATCGSKQKAQ	50
LCSQQVKYLG LKLSKVTRAL REERIQRILA YPHPKTLKQL RXFLGITAF	100
15 RIWIPRYSEI ARPLTLSKE TQKANTHIVR WTPETEVAFQ ALK	143

## (2) INFORMATION FOR SEQ ID NO: 183:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 25 base pairs
  - (B) TYPE: nucleotide
  - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear
- (ii) TYPE DE MOLECULE: ADNc
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:183:

25 GGCCAGGCAT CAGCCCAAGA CTTGA 25

## (2) INFORMATION FOR SEQ ID NO: 184:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 22 base pairs
  - (B) TYPE: nucleotide
  - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear
- (ii) TYPE DE MOLECULE: ADNc
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:184:

35 TGCAAGCTCA TCCCTSRGAC CT 22

206

## (2) INFORMATION FOR SEQ ID NO: 185:

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 23 base pairs
- (B) TYPE: nucleotide
- 5 (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

(ii) TYPE DE MOLECULE: ADNc

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:185:

GACTTGAGCC AGTCCTCATA CCT

23

10

## (2) INFORMATION FOR SEQ ID NO: 186:

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 22 base pairs
- (B) TYPE: nucleotide
- 15 (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

(ii) TYPE DE MOLECULE: ADNc

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:186:

CTTTAGGGCC TGGAAAGCCA CT

22

20

TABLE NO. 5

SEQUENCES GENERATED BY 'PAN-BETTROVIRUS' PCR OF DENSITY GRADIENT FRACTIONS  
 (containing the peak of RT-activity or the corresponding control fraction)

CULTURE	MSRV c-pol	ERVg(v)	PCR artefacts(v)	Total clones
LM7P (I)	18	4	6	26
PLI-1 (II)	0	1	13	23
MS B-CELL LINE (III)	9	2	8	19
CONTROL B-CELL LINE (IV)	0	0	26	26

- I LM7-infected choroid plexus cell culture.
- II MS patient-derived choroid plexus cell culture (PLI-2).
- III MS patient-derived spontaneous B-cell line (immortalized by endogenous EBV strain).
- IV Non-MS control B-cell line.
- V Clones with >90% homology with the GenBank sequence HSERVg are designated ERVg in this study.
- VI PCR artefacts included primer multimers, concatemers, single primer amplifications, etc.

TABLE NO. 6

## DETECTION OF MSRV IN THE CSF OF PATIENTS WITH MULTIPLE SCLEROSIS AND OTHER NEUROLOGICAL DISEASES

Patient <sup>1</sup>	Age/Sex	Diagnosis	MS Type	MS Activity	MS Duration	MS Treatment at Sampling	MSRV ELISA
ITMS1	27 yrs / M	multiple sclerosis	2° progressive	slow progression	5 yrs	corticosteroids	negative
ITMS2	55 yrs / M	multiple sclerosis	1° progressive	slow progression	8 yrs	none	positive
ITMS3	51 yrs / F	multiple sclerosis	1° progressive	slow progression	2 yrs	none	negative
ITMS4	22 yrs ! F	multiple sclerosis	relapsing remitting	In remission	0 yrs	none	positive
ITMS5	27 yrs / F	multiple sclerosis	1° progressive	slow progression	8 yrs	cyclophosphamide	negative
ITMS6	33 yrs / M	multiple sclerosis	2° progressive	slow progression	16 yrs	none (previously cycloph.+ corticost.)	negative
ITMS7	33 yrs / F	multiple sclerosis	2° progressive	slow progression	8 yrs	none	positive
ITMS8	25 yrs / F	multiple sclerosis	relapsing remitting	stable	3 yrs	—	positive
ITMS9	38 yrs / F	multiple sclerosis	2° progressive	slow progression	3 yrs	none	positive
ITMS10	38 yrs / M	multiple sclerosis	2° progressive	slow progression	7 yrs	corticosteroids	negative
OND1	37 yrs / F	cerebellar atrophy	NA <sup>2</sup>	NA	NA	—	negative
OND2	26 yrs / F	viral myelitis	NA	NA	NA	—	negative
OND3	38 yrs / F	viral encephalitis	NA	NA	NA	—	negative
OND4	28 yrs / F	viral encephalitis	NA	NA	NA	—	negative
OND5	64 yrs / M	viral encephalitis	NA	NA	NA	—	negative
OND6	32 yrs / M	Gullain - Barre	NA	NA	NA	—	negative
OND7	54 yrs / F	cerebrovascular	NA	NA	NA	—	negative
OND8	52 yrs / F	hydrocephalus	NA	NA	NA	—	negative
OND9	25 yrs / F	1° cerebral tumour	NA	NA	NA	—	negative
OND10	21 yrs / M	epilepsy	NA	NA	NA	—	negative

<sup>1</sup> CSF samples from patients ITMS1 - OND2 were made available by Prof. P. Ferrando, University Centre for Multiple Sclerosis, Milan, Italy.<sup>2</sup> CSF samples from patients OND3 - OND10 were made available by Profs. J. Peltat and J. Paret, Dept. of Neurology, University Hospital, Grenoble, France.<sup>2</sup> NA = Not Applicable

**CLAIMS**

1. Nucleic material, in the isolated or purified state, comprising a nucleotide sequence selected from the group including sequences SEQ ID NO:93, SEQ ID NO:94, their complementary sequences and their equivalent sequences, in particular nucleotide sequences displaying, for any succession of 100 contiguous monomers, at least 50% and preferably at least 60% homology with said sequence SEQ ID NO:93, SEQ ID NO:94 and their complementary sequences, excluding HSERV-9 sequence.

2. Nucleic material of claim 1, nucleotide sequence of which is selected from the group including sequences SEQ ID NO:93, SEQ ID NO:94, their complementary sequences and their equivalent sequences, in particular nucleotide sequences displaying, for any succession of 100 contiguous monomers, at least 70% and preferably at least 80% homology with said sequence SEQ ID NO:93, SEQ ID NO:94 and their complementary sequences.

3. Nucleic material, in the isolated or purified state, coding for any polypeptide displaying, for any contiguous succession of at least 30 amino acids, at least 50%, preferably at least 60 %, and most preferably at least 70% homology with a peptide sequence encoded by any nucleotide sequence selected from the group including SEQ ID NO:93, SEQ ID NO:94 and their complementary sequence.

4. Nucleic material, in the isolated or purified state, of retroviral type, comprising a nucleotide sequence identical or equivalent to at least part of the pol gene of an isolated retrovirus associated with multiple sclerosis or rheumatoid arthritis.

5. Nucleic material as claimed in claim 4, said nucleotide sequence being 80 % homologous to said at least part of the pol gene.

6. Nucleic material comprising a nucleotide sequence identical or equivalent to at least part of the pol gene of an isolated virus encoding a reverse transcriptase comprising an enzymatic site comprised 5 between the amino acid domains LPQG and YXDD, said virus having a phylogenetic distance with HSERV-9 of  $0.063 \pm 0.1$ , and preferably  $0.063 \pm 0.05$ .

7. Nucleotide fragment comprising a nucleotide sequence selected from the group including SEQ ID NO:93, 10 SEQ ID NO: 94, their complementary sequences and their equivalent sequences, in particular nucleotide sequences displaying, for any succession of 100 contiguous monomers, at least 50% and preferably at least 60% homology with said sequences and their complementary sequences, said 15 group excluding SEQ ID NO:1, and said nucleotide fragment not comprising nor consisting of the sequence HSERV-9.

8. Nucleotide fragment of claim 7, nucleotide sequence of which is selected from the group including SEQ ID NO:93, SEQ ID NO: 94, their complementary sequences and 20 their equivalent sequences, in particular nucleotide sequences displaying, for any succession of 100 contiguous monomers, at least 70% and preferably at least 80% homology with said sequences and their complementary sequences.

25 9. Nucleotide fragment comprising a coding nucleotide sequence which is at least partially identical to a nucleotide sequence selected from the group including :

SEQ ID NO:93, SEQ ID NO:94; their complementary 30 sequences ; their equivalent sequences, in particular homologous to SEQ ID NO:93, SEQ ID NO:94;

sequences encoding at least part of the peptide sequence defined by SEQ ID NO:95;

35 sequences encoding at least part of a peptide sequence equivalent, in particular homologous to SEQ ID NO:95, which is capable of being recognized by sera of

patients infected with the MSRV-1 virus, or in whom the MSRV-1 virus has been reactivated.

10. Nucleic acid probe for the detection of a virus associated with multiple sclerosis or rheumatoid 5 arthritis, characterized in that it is capable of hybridizing specifically with any fragment according to any one of claim 7 to 9.

11. Probe as claimed in claim 10, consisting of between 10 and 1,000 monomers.

10 12. Primer for the amplification by polymerization of an RNA or a DNA of a viral material associated with multiple sclerosis or rheumatoid arthritis, comprising a nucleotide sequence identical or equivalent to at least one portion of the nucleotide 15 sequence of a fragment as claimed in any one of claims 7 to 9, in particular a nucleotide sequence displaying, for any succession of at least 10 contiguous monomers, preferably 15 contiguous monomers, more preferably 18 contiguous monomers and even most preferably 20 contiguous 20 monomers, at least 70% homology with at least the said portion of the said fragment.

13. Primer as claimed in Claim 12, comprising a sequence selected from the group consisting of SEQ ID NO: 99 to SEQ ID NO: 111.

25 14. Polypeptide encoded by any open reading frame belonging to a nucleotide fragment as claimed in any one of claims 7 to 9.

15. Polypeptide of claim 14, characterized in that the open reading frame encoding it, is comprised, in 30 the 5'-3' direction, between nucleotide 18 and nucleotide 2304 of SEQ ID NO:93.

16. Polypeptide according to claim 15, comprising a peptide sequence at least partially identical to SEQ ID NO: 95.

35 17. Polypeptide, comprising a peptide sequence at least partially identical to SEQ ID NO: 96.

18. Polypeptide of claim 17 exhibiting an enzymatic activity consisting of proteolytic activity.

19. Polypeptide, characterized in that the open reading frame encoding it begins, in the 5'-3' direction, 5 at nucleotide 18 and ends at nucleotide 340 of SEQ ID NO:93.

20. Polypeptide exhibiting an inhibitory activity on the proteolytic activity of polypeptide of claim 18.

10 21. Polypeptide, comprising a peptide sequence identical or equivalent to SEQ ID NO: 97.

22. Polypeptide of claim 21, comprising a peptide sequence identical or equivalent to SEQ ID NO: 98.

15 23. Polypeptide, characterized in that the open reading frame encoding it begins, in the 5'-3' direction, at nucleotide 341 and ends at nucleotide 2304 of SEQ ID NO:93.

24. Polypeptide, characterized in that the open reading frame encoding it begins, in the 5'-3' direction, 20 at nucleotide 1858 and ends at nucleotide 2304 of SEQ ID NO:93.

25. Polypeptide of claim 21 or 23, exhibiting a reverse transcriptase activity.

26. Polypeptide of claim 22 or 24, exhibiting a 25 ribonuclease H activity.

27. Polypeptide exhibiting an inhibitory activity on the reverse transcriptase activity of polypeptide of claim 25.

28. Polypeptide having an inhibitory activity 30 on the ribonuclease H activity of polypeptide of claim 26.

29. Antigenic polypeptide recognized from the sera of patients infected with the MSRV-1 virus, and/or in whom the MSRV-1 virus has been reactivated, characterized in that its peptide sequence is at least partially 35 identical or is equivalent to a sequence selected from the group consisting of SEQ ID NO:95, and fragments thereof,

in particular SEQ ID NO:96, SEQ ID NO:97 and SEQ ID NO: 98.

30. Mono- or polyclonal antibody directed against the MSRV-1 virus, characterized in that it is obtained by the immunological reaction of a human or animal body or cells to an immunogenic agent consisting of an antigenic polypeptide of claim 29.

31. Reagent for detection of the MSRV-1 virus, or of an exposure to the said virus, characterized in that it comprises at least one reactive substance selected from the group consisting of a probe as claimed in claim 10 or 11 ; a polypeptide as claimed in any one of claims 14 to 29 ; or an antibody as claimed in claim 30.

32. Diagnostic, prophylactic or therapeutic composition, in particular for inhibiting the expression of a virus associated with multiple sclerosis or rheumatoid arthritis, and/or the enzymatic activity of the proteins of said virus, said composition comprising a nucleotide fragment of any one of claims 7 to 9.

20 33. Diagnostic, prophylactic or therapeutic composition comprising a polypeptide of any one of claims 14 to 29, or an antibody of claim 30.

34. Process for detecting a virus associated with multiple sclerosis or rheumatoid arthritis, in a biological sample, characterized in that an RNA and/or a DNA presumed to belong or originating from said virus, or their complementary RNA and/or DNA, is/are brought into contact with a nucleotide fragment according to any one of claim 7 to 9.

30 35. Process for detecting the presence or exposure to a virus associated with multiple sclerosis or rheumatoid arthritis, in a biological sample, wherein said sample is brought into contact with a polypeptide, according to any one of claim 14 to 29, or an antibody of 35 claim 30.

1/69

## FIG. 1

Consensus            GTTIAAGGGAT A NO CCT C ATC TCT T IGG TCA GGT ACT GGC C CAAG ATC TAG        50  
 Consensus            GGC ACT T CTC AGG T OCAG S N AC T CT G T YCC T TCAG        85

## SEQ ID NO 3 (POL MSRV-1B)

Consensus            GTTCAGGGAT AG 0 0 0 0 C ATC TATT IGG OCA GGC ACT IAG CT CA AACT CT G A        50  
 Consensus            GOCAGT T CTC ATA OCT IGG AC AY T CT YG TOC T T CCGT        85

## SEQ ID NO 4 (POL MSRV-1B)

Consensus            GTTCAR RGAT AG 0 0 0 0 C ATC TATT IGG OOW RGYAT IAG CC CAAG ACT CT G A        50  
 Consensus            GYCAAT T CTC ATA OCT IGG AC AC T CT I G TOC T T YRG        85

## SEQ ID NO 5 (POL MSRV-1B)

Consensus            GTTCAGGGAT AGC 1 0 0 C ATC TATT IGG OCT GGC ATT A ACC CGAG ACT TA A        50  
 Consensus            GOCAGT T CTC ATA OCT IGG AC AC T CT I G TOC T T IGG        85

## SEQ ID NO 6 (POL MSRV-1B)

Consensus            GIGT I GOCAC AGGG GT TTA R RGAT A NC Y CAT C IM T TIG GY W R G Y AYT  
 Consensus            RRCYCRAKAY YIRRG YCAVT TCT YAKR SY RGS NAY TCT B KYO CT TYR GT  
 Consensus            ACAT GGAT GA C

## SEQ ID NO 7 (POL MSRV-1B)

2/69

## FIG. 2

## CONSENSUS A

SEQ ID NO 3

GTTCAGGGATAGCCC CCATCTATTGGCCA GGCACTAGCTCAATA CTTGAGCCAGTCTC 60  
 V . G . P S S L W S G T G P R S R P L L  
 F R D S P H L F G Q V L A Q D L G H F S  
 L G I A L I S L V R Y W P K I . A T S Q

85

AGGTCCAGGCACTCT GTTCCTTCAG  
 R S R H S V P . S  
 G P G T L F L Q  
 V Q A L C S F

## CONSENSUS B

SEQ ID NO 4

GTTCAGGGATAGCCC CCATCTATTGGCCA GGCACTAGCTCAATA CTTGAGCCAGTCTC 60  
 V Q G . P P S I W P G T S S I L E P V L  
 F R D S P H L F G Q A L A Q Y L S Q F S  
 S G I A P I Y L A R H . L N T . A S S H

86

ATACCTGGACACTCT TGTCCCTCGGT  
 I P G H S C P S  
 Y L D T L V L R  
 T W T L L S F G

## CONSENSUS C

SEQ ID NO 5

GTTCAGGGATAGCCC CCATCTATTGGCCA GGCATTAGCCCAAGA CTTGAGTCATACTC 60  
 V Q G . P P S I W P G I S P R L E S I L  
 F R D S P H L F G Q A L A Q D L S Q F S  
 S G I A P I Y L A R H . P K T . V N S H

85

ATACCTGGACACTCT TGTCCCTTCAG  
 I P G H S C P S  
 Y L D T L V L Q  
 T W T L L S F

## CONSENSUS D

SEQ ID NO 6

GTTCAGGGATAGCTC CCATCTATTGGCCT GGCATTAACCCGAGA CTTAAGCCAGTCTC 60  
 V Q G . L P S I W P G I N P R L K P V L  
 F R D S S H L F G L A L T R D L S Q F S  
 S G I A P I Y L A W H . P E T . A S S H

85

ATACGTGGACACTCT TGTCCCTTG  
 I R G H S C P L  
 Y V D T L V L W  
 T W T L L S F

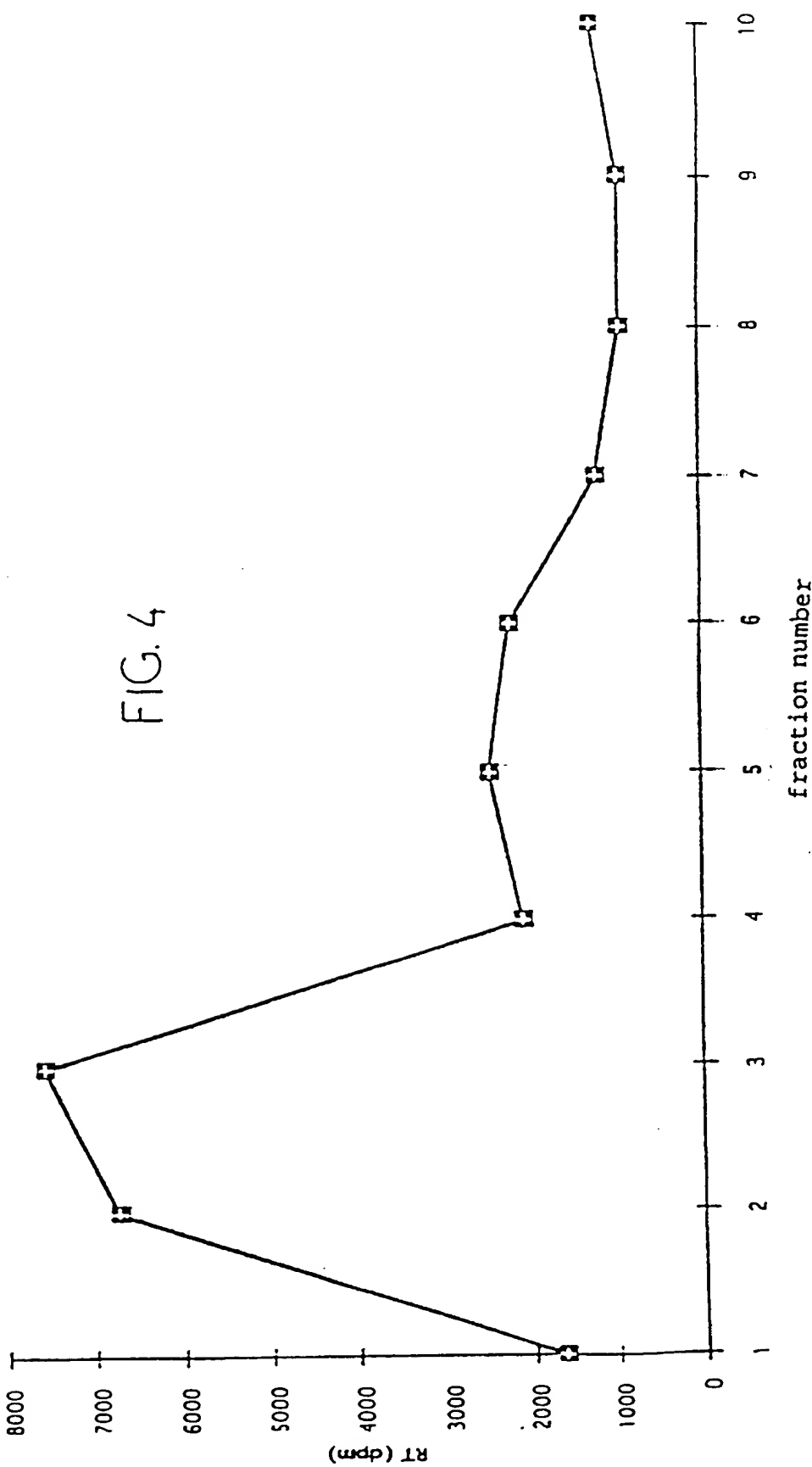
3/69

## FIG. 3

Consensus	TTCGGATOCAG TGYTGOCACA CGGCGCTGAA GCGTATOGGG TCCAGTTCGC	50
Consensus	CGATGCCGCC TATAGCCTCT ACGTGGATGA CCTSCIGAAG CTTGAG	96

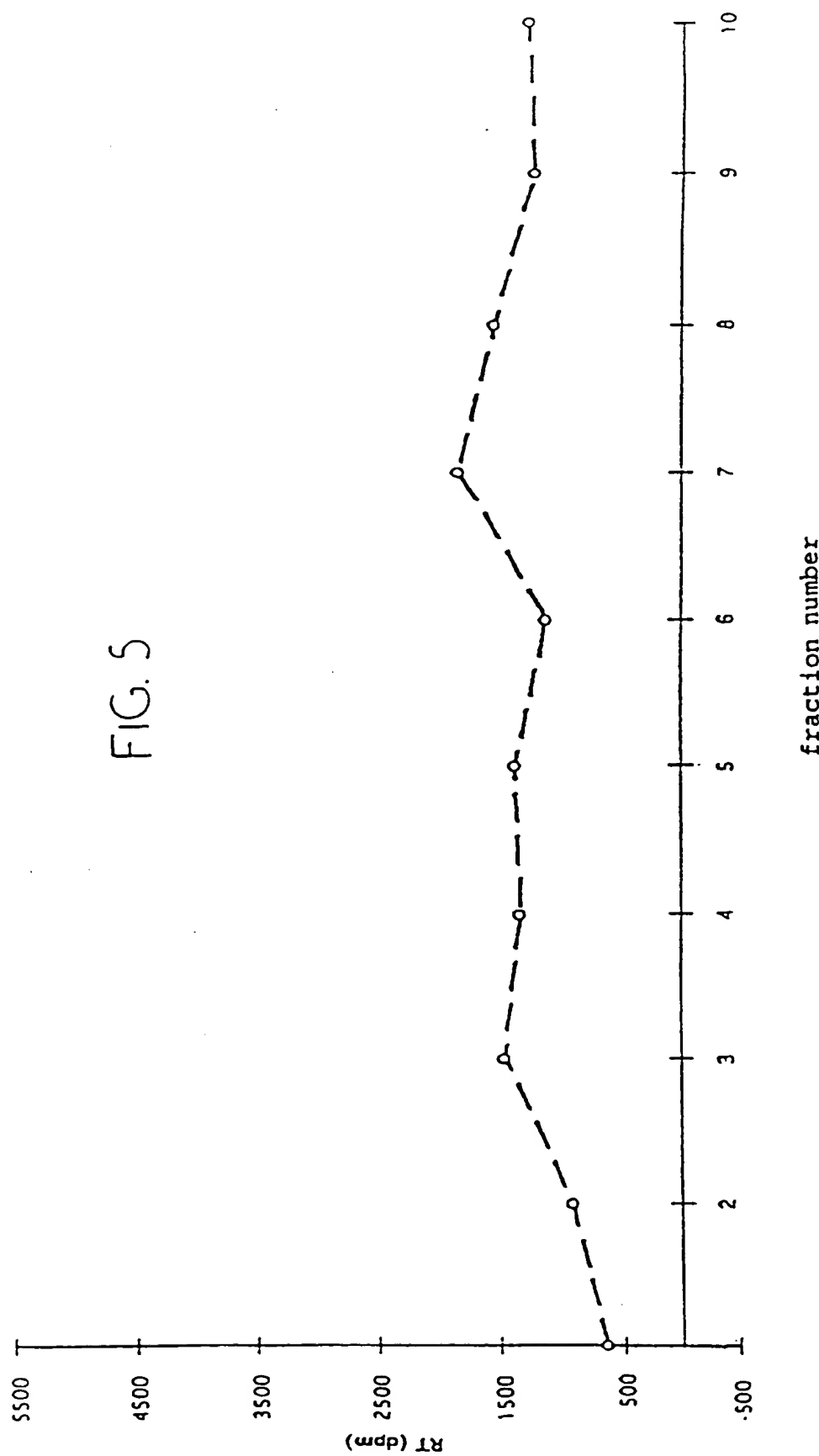
SEQ ID NO 11

4/69



5/69

FIG. 5



6/69

## FIG. 6

CAAGCCACCC AAGAACTCTT AAATTTCCTC ACTAACCTGTG CCTACAAAGT	50
TTCCAAACCA AAGGCTCAGC TCTGCTCACA CGAGATTAGA TACTTAGGGT	100
TAAAATTATC CAAAGGCACC AGGGGCCCTCA GTGAGGAACG TATCCAGCCT	150
ATACTGGTT ATCCTCATCC CAAAACCCCTA AAGCAACTAA GAGGGTTCCCT	200
TAGCATGATC AGGTTTCTGC CGAAAACAAG ATTCCAGGT ACAACAAAAA	250
TAGCCAGACC ATTATATACA CTAAATTAGG AAACTCAGAA AGCCAATAAC	300
TATTTAGTAA GATGGACACC TAAACAGAAG GCTTTCAGG CCTTAAAGAA	350
GGCCTTAACC CAAGCCCCAG TGTTCAGCTT CCCAACAGGG CAAGATTTTT	400
CTTTATATGG CACAGAAAAA ACAGGAATCG CTCAGGAGT CCTTACACAG	450
GTCGGAGGGA TGAGCTTGCA ACCCGTGGCA TACCTGAATA AGGAAATTGA	500
TGTAGTGGCA AAGGGTTGGC CTCATNGTT ATGGGTAATG GNGGCAGTAG	550
CAGTCINAGT ATCTGAAGCA GTTAAAATAA TACAGGGAAG AGATCTINCT	600
GIGTGGACAT CTCATGATGT GAAOGGCATA CTCACTGCTA AAGGAGACTT	650
GIGGTGICA GACAACCATT TACTTAAATA TCAGGCTCTA TTACTTGAAG	700
AGOCAGTGCT GNGACTGOGC ACTTGTGCAA CTCCTAAACC C	741

SEQ ID NO 9 (PSJ 17)

7/69

TCAGGGATAGCCCCATCTATTTGCCAGGCATTAGCCCAAGACTTGAGTC  
AATTCTCATACCTGGACACTCTTGTCTTCAGTACATGGATGATTTACTTT  
TAGTCGCCCGTTCAGAAACCTTGTGCCATCAAGCCACCCAAGAACTCTTAA  
CTTCTCACTACCTGTGGCTACAAGGTTCCAAACCAAAGGCTGGCTCT  
GCTCACAGGAGATTAGATACTNAGGGCTAAAATTATCAAAGGCACCAGG  
GCCCTCAGTGAGGAACGTATCCAGCCTATACTGGCTTATCCTCATCCAAA  
ACCTAAAGCAACTAACAGAGGGTCTTGGCATAACAGGTTCTGCCGAAA  
ACAGATTCCCAGGTACASCCCAATAGCCAGACCATTATACACTAATTA  
NGGAAACTCAGAAAGCCAATACCTATTTAGTAAGATGGACACCTACAGAA  
GTGGCTTCCAGGCCCTAAAGAAGGCCCTAACCCAGGCCAGTGTTCAGC  
TTGCCAACAGGGCAAGATTTCTTATATGCCACAGAAAAAACAGGAAT  
AGCTCTAGGAGTCCTTACGCAGGTCTCAGGGATGAGCTTGCAACCGTGGT  
ATACCTGAGTAAGGAAATTGATGTAGTGGCAAAGGGTT

SEQ ID NO 8 (MO03-PO04)

FIG. 7

8/69

290

TTC AAG GCA  
F K G>  
a a a

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FIG. 9

SEQ ID NO 1 (MSRV-1 pol.)

10/69

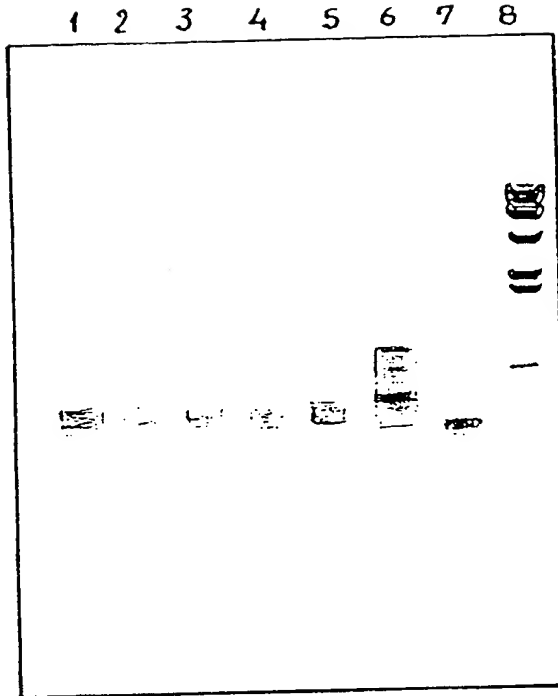
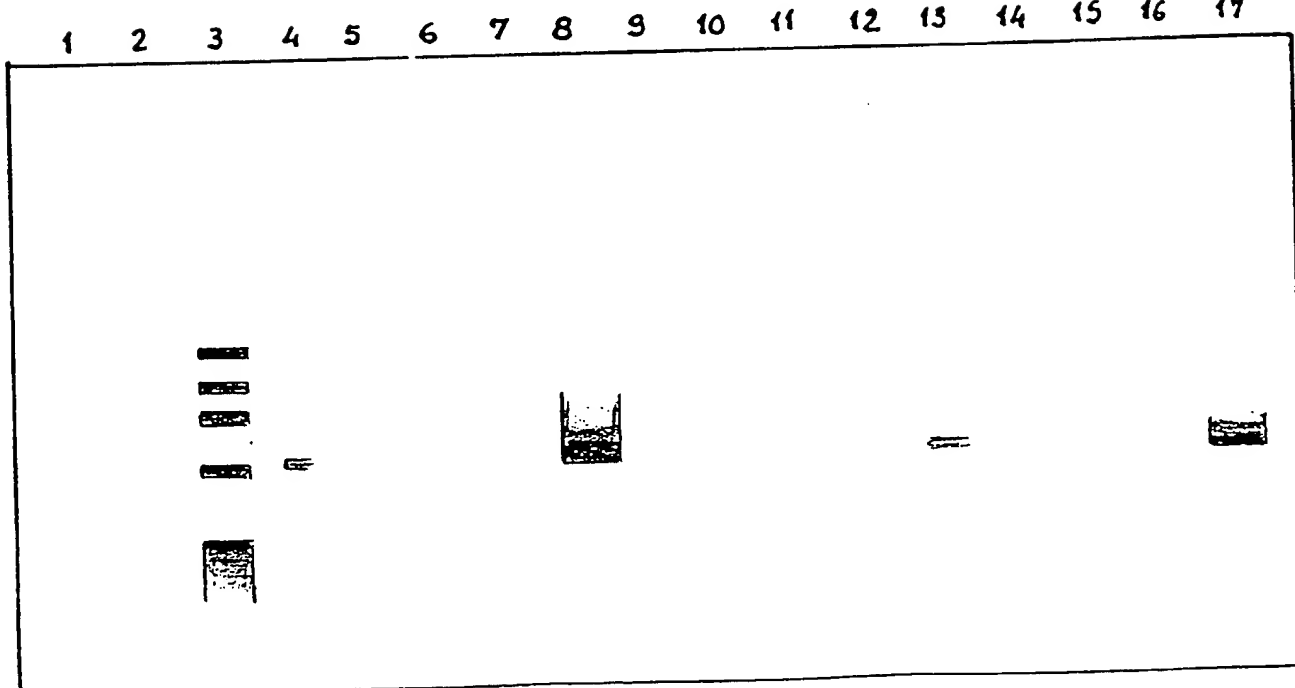


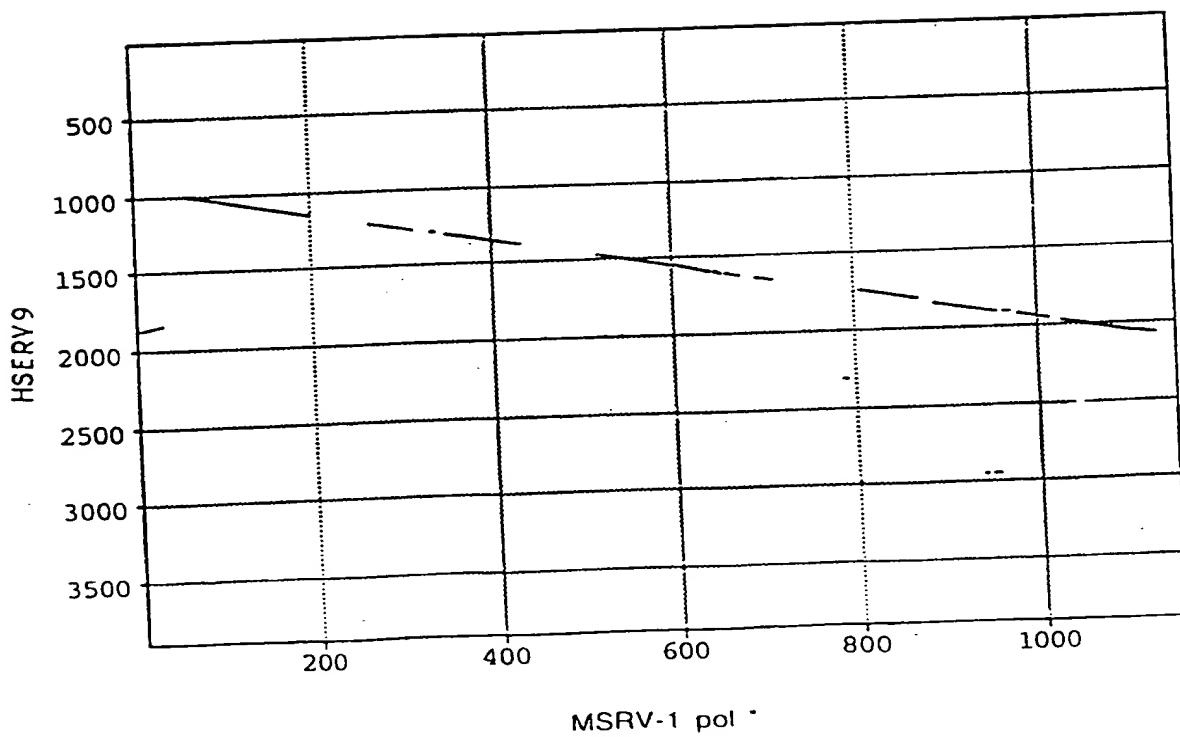
FIG. 10

FIG. 11



11/69

FIG. 12



12/69

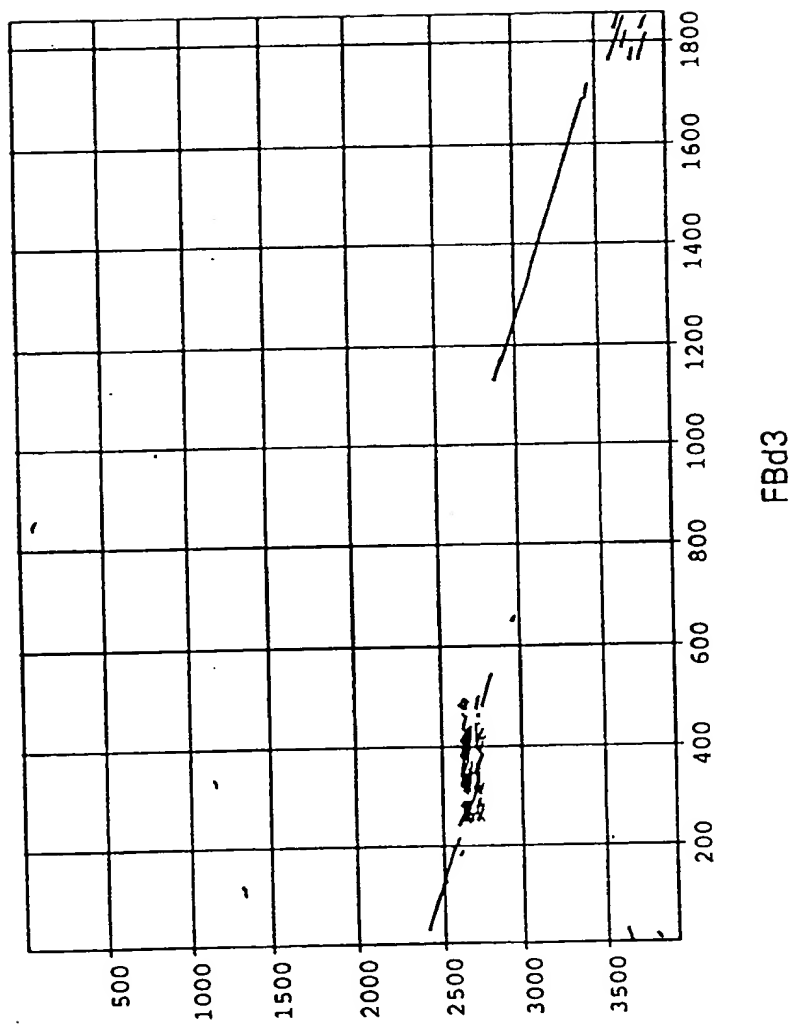
## FIG. 13

SEQ ID NO 46 (FBd3)

GTGCTGATTGGTGTATTAACAATCCTTATCTAATCCGAAATGCCCATGTTG  
CAATATGGAAAGAAAGGGAGTCCTAACCTCTGGGGGAACCCCCATTAAA  
TACCACAAGTAAATCATGGAGTTATTGCACACAGTCAAAAACTCAAGGA  
GGTGGAAAGTCTTACACTGCCAAAGCCATCAGAAAAGGGAAGAGGGGAGAA  
GAGCAGCATAAGTGGCTACAGAGGCAAGGAAGACTAGCAGAAAGGAAA  
GAGAGAAAGAGACAGAAAGTCAGAGAGAGAGAGACAGAGAGTCAGAGAGAAGGAA  
CAAAGAGGGAGTCAGAGAGAGAGAGACAGAGAGTCAGAGAGAAGGAA  
AGAGAGAGAGGAAGAGACAAAGAATGAATCAAACAGAGAGACAGAAAGT  
CAGAGAGAGAGAGAGAGAGAGACAGAGAGAGAGAGAGAGAGTCAGAA  
AAAGAGAGACCAAAGAAGAAGTCCAAAGAGAAAGAAAGAGAGATGGAAG  
TAGTAAAGGAAAAACAGTGTACCTTATTCTTAAAAGCCGGGTAAATT  
AAAACCTATAATTGATAACTGAAGGTCTCTGTAAACCTGTAAACACTCC  
AATACCACCTTGTGTCAAGTGTAAACAAGGGCGTAGCCAAAAGCACTG  
AGGCCACTAACAAACCCATAGCCTCCTATCAAATTCTTAACCCAGCAGG  
TTCTAACAGGGGATCTAAATCTTAATTACCATACAATGGTCCAAC  
CAGACTTAGGAGGAATTCCCTTCAGGACGGGAAGATAGATGCTTCTCCC  
GGCGATTAAGGGAGAAAGACACAATGGGTATTCAAGTAAAGTGCCAAGGGGA  
ACACTTGTAGAAGCAAAGTTAGAAAATTGCCAAATAATTGGTTGCTCAA  
GAGTTGTTGCACTCAGCCAAACCTTGAAGTACTTGCAAGAATCAGAAAGGA  
GCCATCTATACCAATTCTAAGTTAAATATGGACTGAAGGGAGTTTATTAAAT  
ACCAAAGAGAAATTAAAATCCCAAACCTTATAAGGTTTCAACCAAAGTAA  
AGTTGCTAAAAGTTAACAGCGTAACATGTATTATCCTACTACCACACACT  
CTCAAAGGATTCTCAGACAGTTGCAAGAAATAATGATATCTATCCTTAC  
TCTACAATCCCAAATAGACTCTTGGCAGCAGTGAAGTCTCCAAAACCGTCA  
AGGCCTAGACCTCTCACTGCTGAGAAAGGAGGACTCTGCACCTTCTTAAG  
GGAAGAGTGTCTTACACTAACAGTCAGGGATAGTATGAGATGCTGC  
CCGGCATTACAGAAAAAGGCTCTGAAATCAGACAACGCCCTTCAAATT  
CTATACCAACCTCTGGAGTTGGCAACATGGTTCTCCCTTATGTCCC  
ATGGCTGCCATCTGCTATTACTCGCCTTGGGCCCTGTATTAAACCTCC  
TTGTCAAATTGTTCTCTAGGATCGAGGCCATCAAGCTACAGATGGTCTT  
ACAAATGGAACCCCAAATGAGCTCAACTATCAACTTCTACTGAGGACCCCT  
AGACCAACCCCTGGCCCTTCACTGGCCTAAAGAGTCCCCTGGAGGA  
CACTACCACTGCAGGGCCCCATCTTGGCCCTATCCAGAAGGAAGTAGCTA  
GAGCAGTCATTGCCAATTCCCAAGAGCAGCTGGACTCTGGTGGGGACTTG  
GGGGATTGAGAGGTGAAGCCAGCTGGACTCTGGTGGGGGGACTTG  
GAGAACTTTGTGTCTAGCTAAAGGATTGTAATGCAACAATCAGTGCCT  
GTGTCTAGCTAAAGGATTGTAATACACCAATCAGCAC

13/69

FIG. 14



HSERY9

14/69

FIG. 15

SEQ ID NO 51 ( t pol)

GGCTGCTAAAGGAGACTTGTGGTTGTCAGACAATGCCTACTTAGGTACCA  
GGCCTTATTACTTGAGGGACTGGTGCTTCAGATGCGACTTGTGCAGCTCT  
TAACCCAAACTTATGCTGCCAGAAGGATCTTTAGAGGTCCCCCTAGCCA  
ACCCTGACCTAACCTATATATATACTGATGGAAGTTCGTTGTAGAAAAG  
GGATTACAAAGGGNAGGATATNCCATAGGTTAGTGATAAAGCAGTACTTG  
AAAGTAAGCCTCTCCCCCAGGGACCAGCGCCCCCGTTAGCAGAACTAGT  
GGCACTGACCCCGAGCCTAGAACCTGGAAAGGGAGGAGGATAAATGTGT  
ATACAGATAGCAAGTATGCTTATCTAACCGAAATGCCATGTTG

15/69

SEQ ID NO 52 (JLBc1)

TCAGGGATAGCCCCATCTATTGGTCAGGCCTGGCCAAAGATCTAGGGA  
CATGCCACTTTAAGAGCCATTCTCAAGTCCAGGTACTCTGGTCTTCGGT  
ATGTGGATGATTACTTTGGCTACCAAGTTCAGTAGCCTCATGCCAGCAGG  
CTACTCTAGATCTCTGAACCTTCTAGCTAATCAAGGGTACAAGGCATCTA  
GGTTGAAGGCCAGCTTGCTACAGCAGGTCAAATATCTAGGCCTAATCT  
TAGCCAGAGGGACCAGGGCACTCAGCAAGGAACAAATACAGCCTATACTG  
GCTTATCCTCACCCCTAACGACATTAAAACAGTTGCGGGGGTCTTGGAAATC  
ACTGGCTTTGGTACTATGGATTCCCAGATACAGCAAGATTGGCAGGCC  
CCTCTATACTGTAATCAAGGAGACTCACGAGGGCAAGTACTCATCTAGTAG  
AATGGGAACTAGGGACAGAAACAGCCTTAAACCTTAAAGCAGGCCCTA  
GTACAATCTCCAGCTTAAGCCTTCCCACAGGACAAAACCTCTTATAC  
ATCACAGAGAGGGCAGAGATAGCTTGGTGTCTTATTCAAGACTCATGGG  
ACTACCCCCACAACCAAGTGGCACACCTAACGAAATTGATGTAGTAGC  
AAAAGGCTGGCCTCACTGTTATGGTAGCTGTGGTGGCTGTCTTAGT  
GTCAGAAGCTATCAAATAACAAGGAAAGGATCTCACTGTCGGACTA  
CTCATGATGTAATGGCATACTAGGTGCCAAAGAAGTTATGGGTATCAGA  
CAACCACCTGCTTAGATACCAGGGACTACTCCTGGAGGATTGGGCTTCAAG  
TGCCTTTTGTCGGCTCAACCCTGCCACTTTCTCCAGAGGATGGAGAG  
CCGCTTGAGCATGCTTGCACAGGTTGAGGCCAGAATTATTCCACCCGA  
GATGATCTTAGAGTACCCCTAGCTAACCTGACCTAACCTATATACCA  
ATGGAAGTTCTTGAGGAAACGGGATATGAAGGGCAGGTTATGTCTAG  
TTAGTGATGTAATCATACTGCAAGTAAGCCTTACCCAGGGCCAGCA  
CTCAGTTAGCAGAACTAGTCACACTAACCTAACCTAGAACTGGGAAAGG  
GAAAAAGAATAATATGTATACAGATAGTAAGTATGCTTATCTAACCTAC  
ATGCCCATGTCGAATATGGAAGGAAGGGAGTTCTAACCCCTGGGGGA  
ACCCCATTAATACCAAGGYAAATCATGGAGTTATTGCACGCAGTGC  
AAAAACTCAAGGAGGTGGCAGTCTACACTGCCGAAGCYATCAAAAGGG  
GAAGGAGAGGGGAGAACAGCAGCATAAGTGGTGGCAGAGGCAGTGAAA  
GACCAGCAGAGAGAAGGAGAGAGAACGTCAACGACAGAAGGAAAGAA  
GAGGAGGAGACAGAGAGGAAGAGACAGAGAGAACAGTCTAACAGAGAG  
AGACAGAGAGAGGAAGAGACAGACAGAACAGAAAGTCCAAGAGAGAAGGAAAGA  
GAGGAAGAGACCAAGGAGTCCNAGAGAGAGAACAGAGATAGAAGTAGTAA  
AGAAAAAAACATTGTACCCATTCTTAAAGCCGGGTATATTAAAACC  
TATAATTGATAATTGAGTTCTGCACCCCTCCAGGGGATYGTGGGAGG  
AAACCCCTCAACCGATATGTGAAAATTGTTGGTCTGCTTATGTCTCAATT  
CCAGCCAATACCCCTTGTCTTATTGTAACGAGGGTGTAGAGCGCAGAC  
AGGGAGACCTGTACAATCCATACCCCTCTAACCTAAATCTAACCCAG  
CAGGTTCTAAAGGGGATCTAAATCTAACCTAACCTAACAGGTC  
AAACCCAGATCTAGGAGGAACCTCCTCAGGACAGGATGATAGATGGTCT  
CCCAGGCATAAGAAAATAAAAGACACATGGCAGCCAGTAAGTGAT  
AAGGGAAACACTAGTAAAGCAGTTAGGAGAAGTGTCTAACCTAACAGGTT  
ACTCCAAATGTGTGAGTTGTTGCACTCAGCCAAATCTAAAGTACTTAC  
AGAATTAGGGAGGAGCCATTACACCAATTCTAACGTTAAATATGGACTGGAT  
GAGGTTTATTAAATAGCGAAGGAGAATTAAATCTAACAGTGTAAACATGCATTATCCTA  
CAACTAAAGTAAATTACTAAAAGCTAACAGTGTAAACATGCATTATCCTA  
CTACAAACACACTCTCANAGGATTCCCTCAGACAGTTACAAGAAATAACAA  
AATCTATCTGTAAGGATAGTAACTACAATCCAAATACATTCTTGGCAG  
CAGTGAACACTC

FIG. 16

16/69

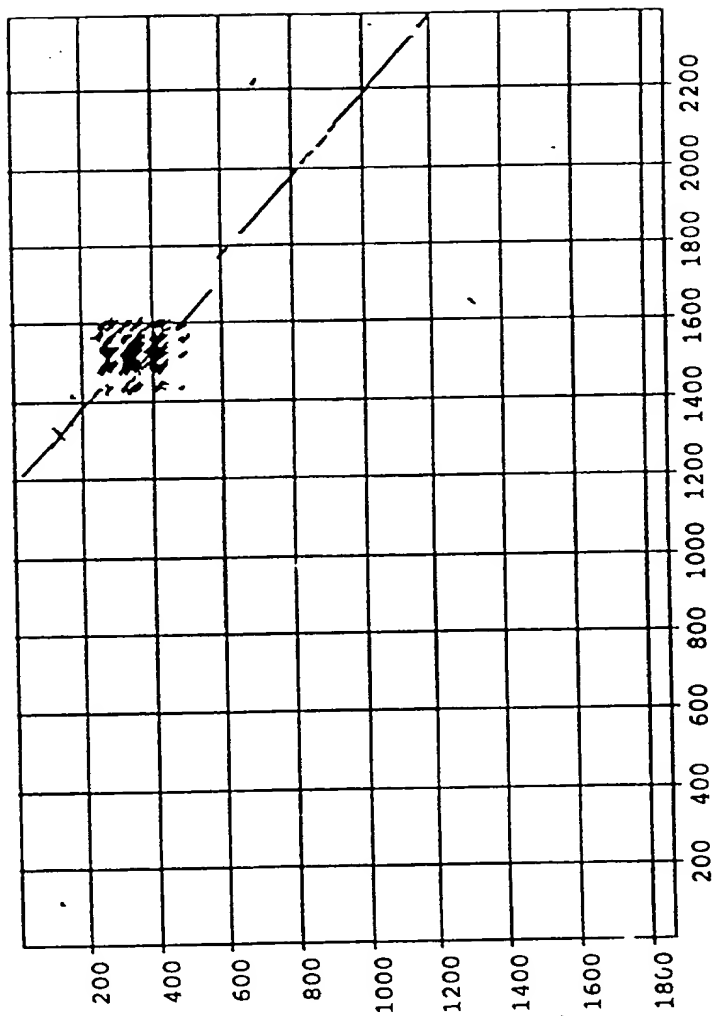
SEQ ID NO 53 (JLBc2)

TCAGGGATAGCCCCCATCTATITGATCAGGCACTAGCCCAAGATCTAGGCC  
ACTTCTGAAGTCCAGGCATTCTAGTCCTCAGTATGTGGATGATTTACTTT  
GGCTACCACTGTTGGAAAGCCTCATGCCAGCAGGCTACTTGAGATCTCTGAA  
CTTTCTAGCTAATCAAGGGTGTATGGCATCTAAATTGAAAGTCCAGCTCTG  
CCTACACAACAAGTCAAATATCTAGGCCTAATCTTAGATAGAAGAACAGGG  
CCCTCAGCAAGGAATGAATAAAGCCTATGCTGGCTATCGGCACCCTAAGA  
CATTAAAACAATTGTGGGGGTTCTTGGAAATCACTGGCTTTGCCGACTAT  
GGATCCCTGGATAGAGTGAGATAGCCAGGCCCTCTATTACTCTTATCAA  
GGAGACCCAGAGGGCAAATACTTATCTAGTATTATGGGNACCAGAGGCAG  
AAAAAGCCTTCAAACCTTAAAGGAGACCTAGTACAAGCTCCAGCTTAA  
GCCTCCCACAGGACAAANCTTCTTTATATGTACAGAGAGAGCAGGAA  
TAGCTCTGGAGTCCTACTCAGACTTTGGACGACCCACGGCCAGTGGC  
RTACCTAAGTAAGGAAATTGATGATGAGTAGCAAAGGCTGGCCTACTGTTT  
ATGGGTAGTTGCGGCTGTGGCAGTCTTACTGTCAAAGGCTATCAAATAAT  
ACAAGGAAAGGATTCACTATCTGGACTACTCATGAGGAAAATGGCATATT  
AGGTGCCAAAGGAAGTTTGGCTATCAGACAACCACTGCTCAGATTCCA  
GGCACTACTGATTGAGAGACCAGTGCTTAAATATGTATGTGTGTGG  
CCCTCAACCCTGCCACTGTTCTCCCAGAAGATGGAGAACCAATGAAGCATT  
ACTGTCAACAAATTAGAGTCCAGAGTTATGCTGCCTGAGAGGATCTTAG  
AAGTCCCCCTAGCTAATCCTGACCTAACCTATATGCTATGGAAGTTCAC  
TTGTGGAGAATGGGATACGAAAAGCACATTATGCCATAGTTAGTGAGGTA  
ACAGTACTTGAAAGTAAGCCTATTCCCCCATGGACCAGAGCCCAGTTAGCA  
GAACTAGTGGCACTTACCCAAAGCCTAGAAACTAGGAAAGGGAAAATAAT  
AAATGTGTATACAGATAGCAAGTATGCTTATCTAACCTACATGCCATGC  
TGCAGTATGAAAGAAAGGGAGTTCTAACCTCTGGGGGAACCCCCATTA  
AATACCACAAGGCAAATCATGGAGTTATGCTATGAGGAAACCTCAA  
GTAGGTGGCAGTTTACACTGCCTGAAGCTATGGGAAGGAGAGAGGAGA  
ACAGCAGCATAAGTGGCTAGCAGAGGAGCGAAAGACTAGCAGAGAGGA  
GAGGTAGGGAAAGACAGAAAGTCAAAGAAAAGAAGTCAAAGACAGACA  
GAGAAAGAGACAGAGGGAGCCAGAGAGAAAAGAGAGAACAGAAAGA  
GACAGAATGTCAAAGAACAGAACAGAGAGAGGGAGCGCCAGAACAGTTAAG  
AAAGTGAGAAAGAGAGATGGAATAGTAAAGAAAAAACAGTGTACCCSTAT  
TCCTTAAAAGCCAGGGTAAATTAAACGTATAATTATAATTGGAAGG  
TCTTCTCCATAACCCATAACATTAAACCTTGTGTCAGTGTAAAC  
AAGAGCATAGCCAAAAGCACTGAGGCCACTGACAACCCATAGCCTTCT  
ATCAAAAATCCTTAACCTGCAAGGTTCTAACAGGGATCTAAATCTCAA  
CTAACACCATAACATGGTCGACCAGACCTAGGAGCGACTCCCCCTCAGG  
ACAGAAGGATGGATGGTTCTCCAGGCCATTAAAGGGAAAGAGACACAAT  
GGGTATTCACTGATAAGGAAACTCTTGAGAAGCAGTTAGGAAGATT  
GCCTAATATTGGTCTGCTCAAATGTGCCAGCTGTTGCACTCAGCTAAC  
CTTAAATTACTTACAGAATTAGGAAGGAGCCATCTACCAATTCTGAGTT  
AAATGAGCTGAACAAGTTCTTAAATAGCAAAGAATCATTGAAATCTCA  
AACTTGCAAAGTTCAACAAAAGTAAAGTTGCTGAAAGTTAGCAGTGTAA  
ACATGTATTATCCTAACCTCTAACATTGTGGAAATCAGACCCCTATCAGTGC  
CCCTCAAAGCTGAAGTCCATCAGCATATGGCCATACAAACTAACCCCTAT  
TTATAGGGTTAGGAATGGCCACTGCTACAGGAATGGGAGTAACAGGTTAT  
CTACTTCATTATCCTATTACCAACACACTCTTAAAGGATTTCAGACAGTTT  
ACAAGAAAATAACAAAATCTATCCTTACTCTNTARTCCAAATAGRTTCTTT  
GGCAGCAGTGAUTCTC

FIG. 17

17/69

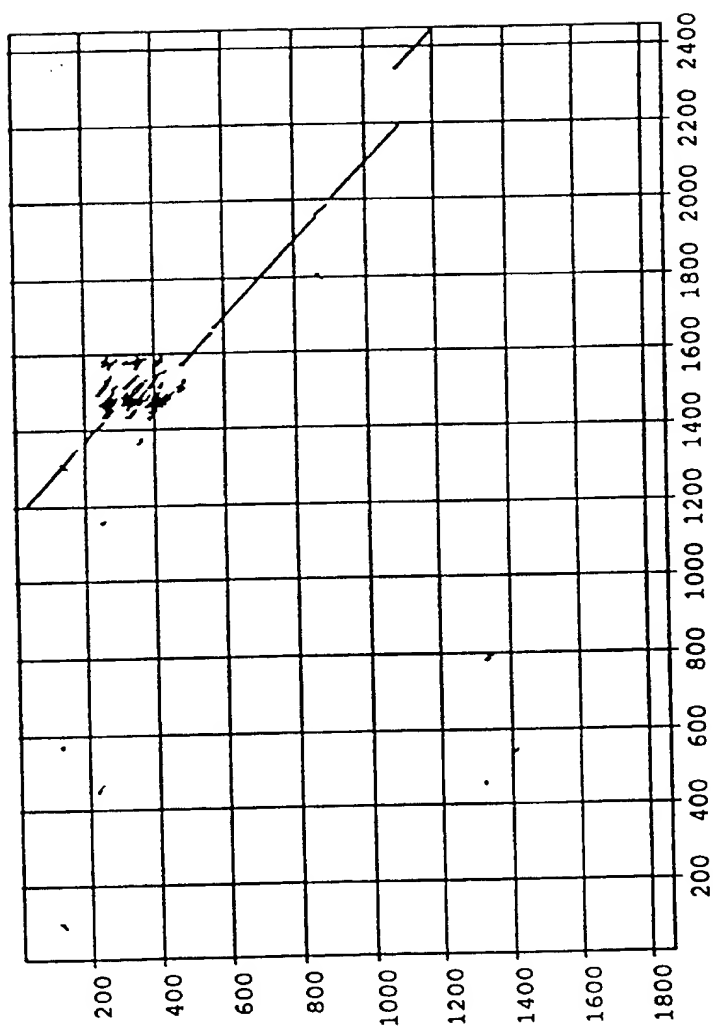
FIG 18



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18/69

FIG 19

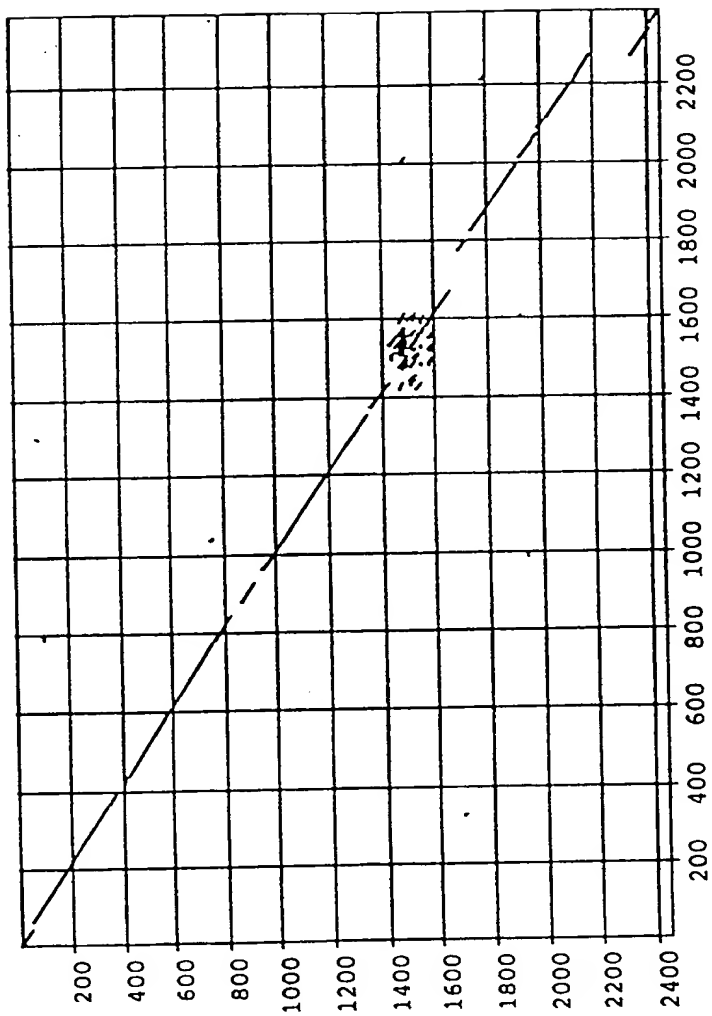


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19/69

FIG 20

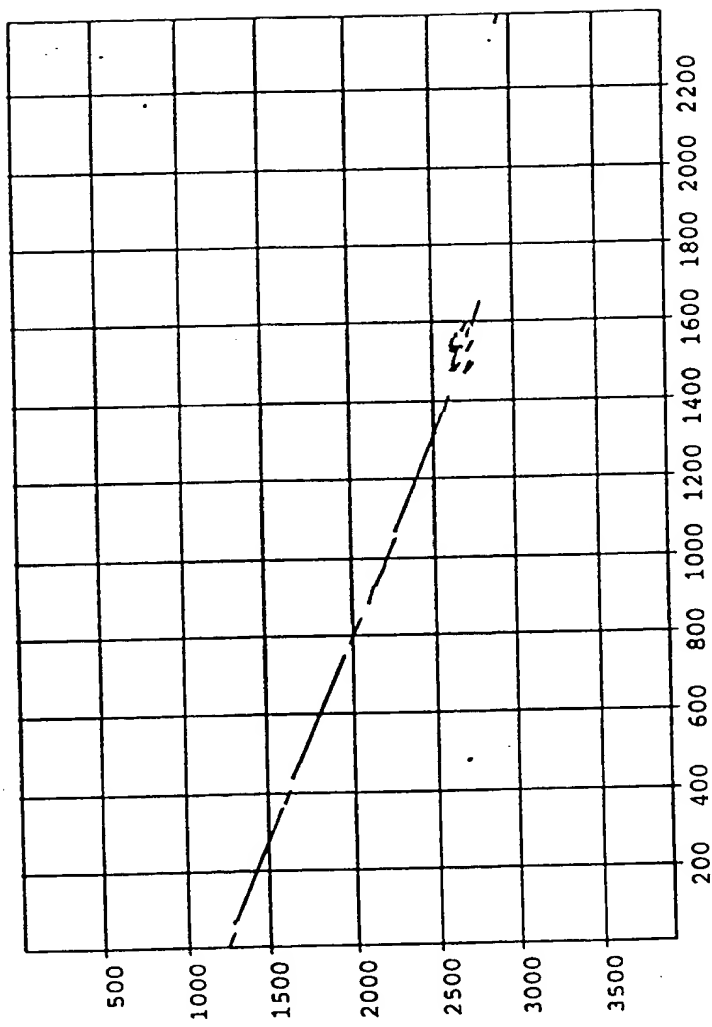


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JLBc1

20/69

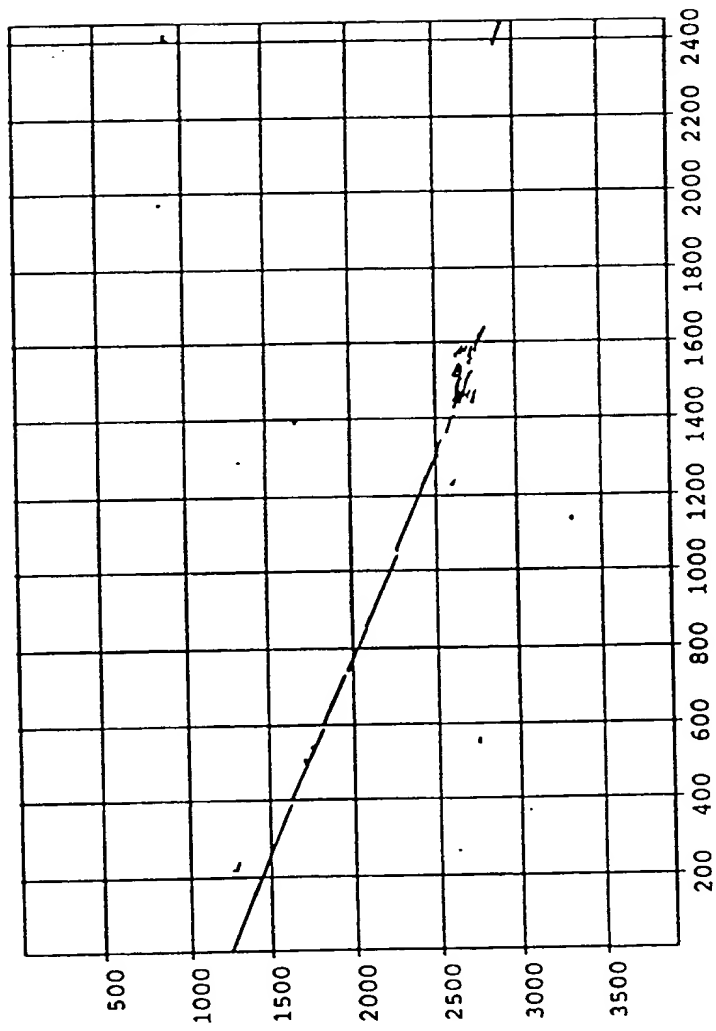
FIG 21



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21/69

FIG 22



JLBc2

HSERY9

*22/69*

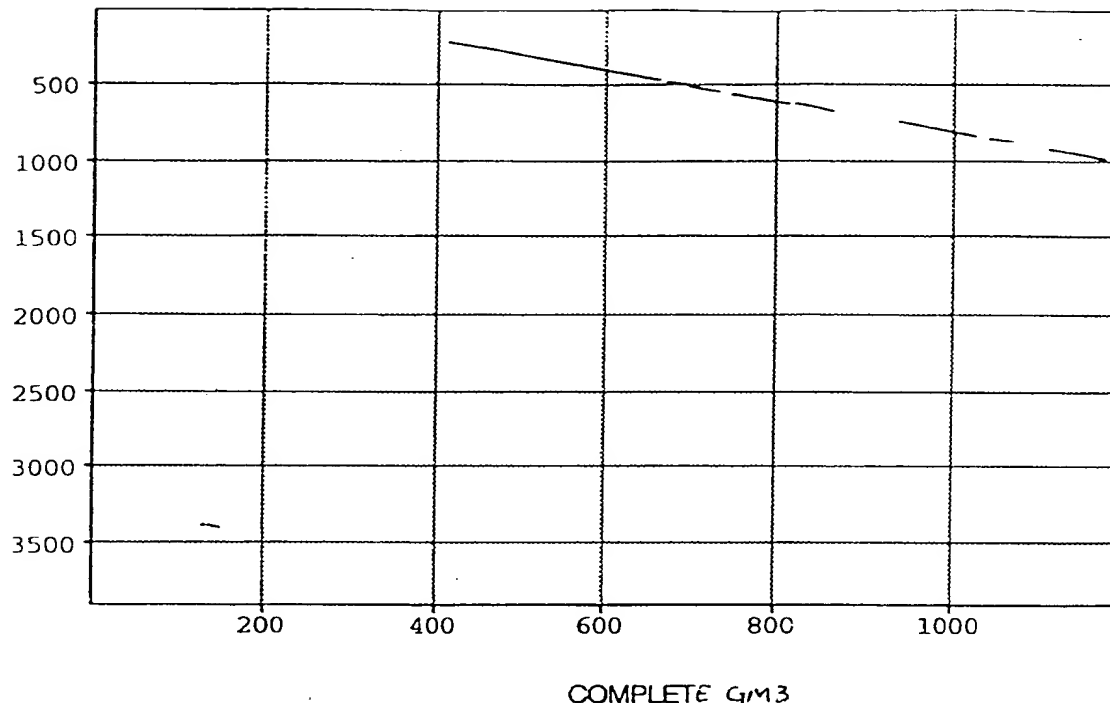
1 TTCCTGAGTT CTTGCACTAA CCTCAAATGA GAGAACTGCC GCCATAACTG CAACCCAAGA  
61 GTTGGCGAT CCCTGGTATC TCAGTCAGGT CAATGACAGG ATGACAACAG AGGAAAGATA  
121 ATGATTCCCC ACAGGCCAGC AGGCAGTCC CAGTGTAGAC CCTCATTAGG ACACAGAAC  
181 AGAACATGGA GATTGGTGC GCAGACATTT GCTAACTTGC GTGCTAGAAG GACTAAGGAA  
241 AACTAGGAAG ATATGAATTA TTCAATGATG TCCACTATAA CACAGGGAA AGGAAGAAA  
301 TCCTACTGCC TTTCTGGAGA GACTAAGGGA GGCATTGAGG AAGCATACCA GGCAAGTGGA  
361 CATTGGAGGC TCTGGAAAAG GGAAAAGTTG GGAAAAGTAT ATGTCTAATA GGGCTTGCTT  
421 CCAGTGTGGT CTACAAGGAC ACTTTAAAAA AGATTGTCCA ATAGAAATAA GCCACCACCT  
481 CGTCCATGCC CCTTATGTCA AGGGAAATCAC TGGAAGGCC ACTGCCCCAG GGGATGAAGG  
541 TCCTCTGAGT CAGAAGCCAC TAACCAGATG ATCCAGCAGC AGGACTGAGG GTGCCCCGGGG  
601 CAAGGCCAG CCCATGCCAT CACCCCTCACA GAGCCCCAGG TATGCTTGAC CATTGAGGGT  
661 CAGAAGGGTA CTGTCTCCTG GACACTGGCG GGCCTTCTCA GTCTTACTTT CCTGTCCTGG  
721 ACAACTGTCC TCCAGATCTG TCACTGTCCG AGGGGTCTA GGACAGCCAG TCACTAGATA  
781 CTTCTCCCAAG CCACTAAGTT GTGACTGGGG AACCTTACTC TTCCACATGC TTTTCTAATT  
841 ATGCCTGAAA GCCCCACTCT CTTGTTAGGG GAGAGACATT CTAGAAAAG CAGGGGCCAT  
901 TATACATGTG AATATAGGAG AAGGAACAAAC TGTTTGTGT CCCCTGCTTG AGGAAGGAAT  
961 TAATCCTGAA GTCCCCGCAA CAGAAGGACA ATATGGACAA GCAAAGAATG CCCGTCTGT  
1021 TCAAGTTAAA CTAAAGGATT CCACCTCCTT TCCCTACCAA AGGCAGTACC CCCTCAGACC  
1081 CGAGACCCAA CAAGAACTCC AAAAGATTGT AAAGGACCTA AAAGCCAAG GCCTAGTAAA  
1141 ACCAAGCAAT AGCCCTTGCA AGACTCCAAT TTTAGGAGTA AGGAAACCCA ACGGAC

SEQ ID NO 56 (GM3)

FIG. 23

23/69

HSERV9



COMPLETE gm3

FIG. 24

24/69

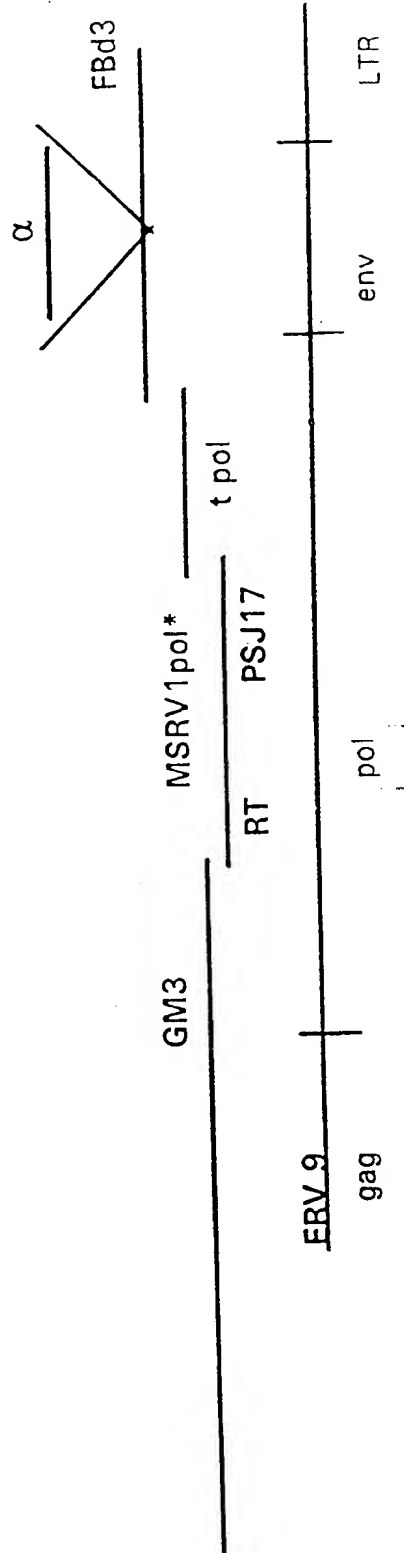
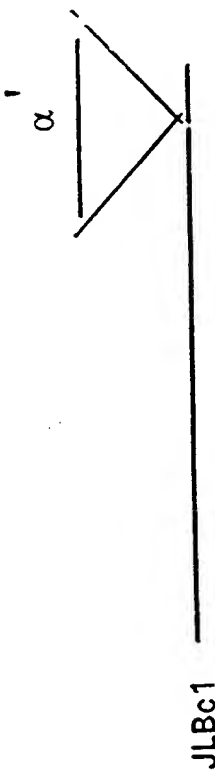


FIG. 25

25/69

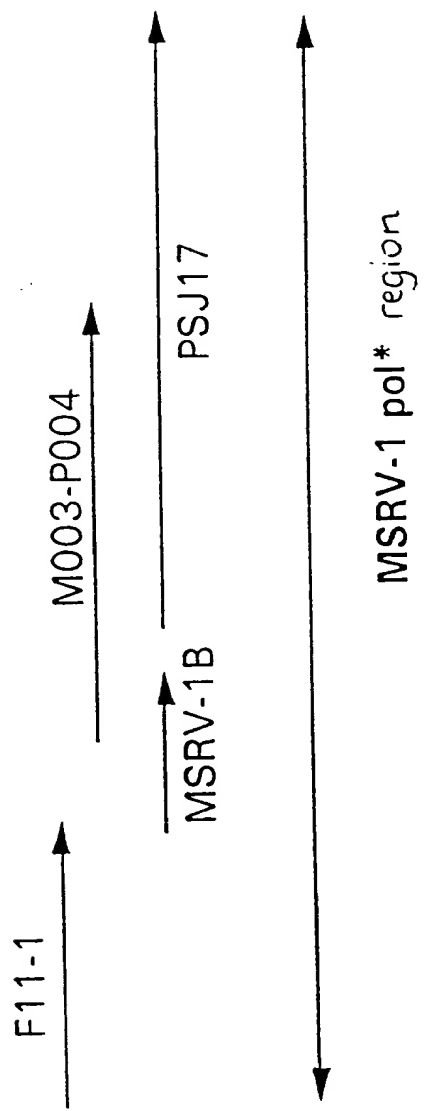


FIG. 26

26/69

27a

SEQ ID NO 57 (POL)

ATG ATC CAG CAG GAC NGA CGG TEC CGG CAA CGG CCA ACC ACC CTC ACA GAG CCC ATC CCT GCC CAT GCC Q A P A H A I T L T E P Q V C L T I E 90  
 M I Q Q D X G C P G C Q D T G C L L D T G C A F S V L L S C P G Q L S S R S V T 100  
  
 CCT CAG AAG GGT NAC TGT CTC CTC CTG GAC ACT GGC TTC GCA GCA CCT TCC TGT CCT GCA CAA CCT TCC TGT CCT GCA TCT GTC ACT 270  
 G C Q K G X C L L D T G C A F S V L L S C P G Q L S S R S V T  
  
 GTC CGA CGG GTC CTA CGA CAG CCA GTC ACT AGA TAC TTC TCC CAG CCA CTA AGT TGT GAC TGG GGA ACT TTA CTC TTC CCA CAT CCT TTT  
 V R G V L G Q P V T R Y F S Q P L S C D W G T L L F P H A F  
  
 CTA ATT ATG CCT GAA AGC CCC ACT CTC TTC TTC GGG AGA GAC ATT CTA GCA AAA CGA CGG GCC ATT ATA CAT GTG AAT ATA GGA GAA CGA 350  
 L I M P E S P T L L G R D I L A K A G A I H V N I G E G  
  
 ACA ACT GTT TGT CCC CTC CTC GAG GAA GAA ATT AAT CCT GAA GTC CGG GCA ACA GAA GAA CAA TAT GCA CAA AAG AAT GCC CGT 450  
 T T V C C P L L E E G I N P E V R A T E G Q Y G Q A K N A R  
  
 CCT GTT CAA GTT AAA CTA AAG GAT TCC ACC TCC TAC CCC TAC CAA AGG CAG TAC CCC CTC AGA CCC GAG ACC CAA CAA GAA CTC CAA AAG 540  
 P V Q V K L K D S T S F P Y Q R Q Y P L R P E T Q Q E L Q K  
  
 ATT GTA AAG GAC CTA AAA GCC CAA CGC CTC GTC AAA CGC AAT AGC CCT TCC AAG ACT CCA ATT TTA CGA GTC AGG AAA CCC AAC CGA 630  
 I V K D L K A Q G L V K P S N S P C K T P I L G V R K P N G  
  
 CAG TCG ACG TTA GTG CGA GAA CTC AGG ATT ATC AAT GAG GGT GTT GTT CCT CTC CTA TAC CCA CCT GTC CCT TAT ACA GTG CTT TCC 720  
 Q W R L V Q E L R I I N E A V V P L Y P A V P N P Y T V L S  
  
 CGA ATA CGA GAA CGA GAG TGG TTT ACA GTC CTC GAC CTT AAG GAT CGC TTT TTC TCC ATC CCT GTC TCA CGT CCT GAC TCT CAA TTC TTG  
 Q I P E E A E W F T V L D A F F C I P V R P D S Q F L 810  
  
 TTT GCC TTT GAA GAT CCT TTC AAC CCA ACG TCT CAA CTC ACC TGG ACT GTT TTA CGG GAT ACC CCC CAT CTA TTT CGC 900  
 F A F E D P L N P T S Q L T W T V L P Q G F R D S P H L F G  
  
 CGC CGA TTA CGC CAA GAC TTG AGT GAA TTC TCA TAC CTC CGC ACT CCT GTC CCT CAG TAC ATG GAT GAT TTA CGT GCC CGT TCA 990  
 Q A L A Q D L S Q F S Y L D T L V L Q M D D L L V A R S  
  
 GAA ACC TTG TGC CAT CGA CGC ACC CGA CGA CTC TTA ACT TTC CTC ACT ACC TGT CGC TAC AGT GTT TCC AAA CGA CGT CGG CTC TGC 1080  
 E T L C H Q A T Q E L T F L T C G Y K V S K P K A R L C

27/69

F.G. 27b

SEQ ID NO 57 (POL)

TCA CAG GAG ATT AGA TAC TTA CGG CTA AAA TCC AAA CGC ACC AGG GCC CTC AGT GAG GAA CGT ATC CAG CCT ATA CTG CCT TAT CCT  
 S Q E I R Y X G L K L S K G T R A L S E E R I Q P I L A Y P 1170

CAT CCC AAA ACC CTA AAG CAA CTA AGA CGG TTC CTT GGC ATA ACA GGT TTC TGC CGA AAA CGG ATT CCC AGG TAC ASC CCA ATA GCC AGA  
 H P K T L K Q L R G F L G I T G F C R K Q I P R Y X P I A R 1260

CCA TTA TAT AGA CTA ATT ANG GAA ACT CAG AAA CCC AAT ACC TAT TTA GTA AGA TGG ACA CCT ACA GAA GTG GCT TTC CAG GCC CTA AAG  
 P L Y T L I X E T Q K A N T Y L V R W T P T E V A F Q A L K 1250

AAG GCC CTA ACC CAA GCC CCA GTG TTC AGC TTG CCA ACA GGG CAA GAT TTT TCT TTA TAT GCC ACA GAA AAA ACA CGG ATA GCT CTA CGA  
 K A L T Q A P V F S L P T G Q D F S L Y A T E K T G I A L G 1440

GTC CTT AGC CAG CTC TCA CGG ATG AGC TTG CAA CCC GTG GTC TAC CTG AGT AAG GAA ATT GAT GTC GTG GCA AAC CGT TCG CCT CAT NGT  
 V L T Q V S G M S L Q P V V Y L S K E I D V V A K G W P H X 1530

TTA TCG GTC ATG CGC GCA GTC TGA TCT GAA CGA GTT AAA ATA ATA CAG CGA AGA GAT CTT NCT GTG TCG ACA TCT CTT GAT  
 L W V M X A V A V X V S E A V K I I Q G R D L X V W T S H D 1630

GTC AAC CGC ATA CTC ACT GCT AAA CGA GAC TTG CGC TCG  
 V N G I L T A K G D L W L S D N H L L X Y Q A L L E E P V 1710

CTG TGA CGC ACT TGT CCA ACT CTT AAA CCC AAA CTT ATG CTG CCC AGA AGG ATC TTT TTA CTT AAN TAT CAG GCT CTA TTA CTT GAA GAG CGA GTG  
 L X L R T C A T L K P R R M L P R I F X E V P L A N P D L 1800

AGC TAT ATA TAT ACT GAT CGA AGT TCG TTT GTC GAA AAG CGA TTA CAA AGG CGA CGA TAT NCC ATA GCT GTT AGT GAT NAA CGA GTT CTT  
 N N Y I Y T D G S S F V E K G L Q R X G Y X I G V S D K A V L 1890

GAA AGT AAG CCT CTT CCC CCC CAG CGA CGA CGG CGG CGG TTA CGA GAA CTA GTG CGA CGG ACC CGG TTA GAA CTT TCG AAA CGC  
 E S K P L P P Q G P A P L A E L V A L T P R A L E L W K G 1980

ACG AGC ATA AAT GTG TAT AGA GAT AGC AAG TAT CCT TAT CTA ATC CGA AAT GCC CAT GTT GTT CTA ATC CGA AAT GCC CAT GTT CGA  
 R R I N V Y T D S K Y A Y L I R N A H V V Y L I R N A H V A 2070

ATA TCG AAA GAA AGG GAG TTC CTA ACC TCT CGG CGA ACC CCC ATT AAA TAC CAC AAG TTA ATC ATG GAG TTA TIG CAC ACA GTG CGA AAA  
 I W K E R E F L T S G G T P I K Y H K L I M E L L H T V Q K 2160

28/69

SEQ ID NO 57 (POL)

CTC AAG GAG GTC GAA GTC TTA CAC TCC CAA ACC CAT CAG AAA AGG GAA ACC GCA GAA GAG CAG CCG CAT AAC TOG CTA CAG AGG CAA GCA AAC 2250  
L K E V L H C Q S H Q K R E R G E E Q H K W L Q R Q G K  
ACT AGC AGA AAG GAA AGA GAG ACA GAA GAG AGA AGT CAG AGA GAG AGA GAG GAA GAG ACA GAG CAC AAA GAG GCA GTC AGA GAG AGA GAG 2340  
T S R K E R E K E T E S . Q R E R E E T E H K E G V R E R E  
AGA CAG AGA GTC AGA GAG AAG GAA AGA GAG AGA GCA AGA GAC AAA GAA GGA TGA  
R Q R V R E K E R E R G R D K E .

FIG. 27c

*29/69*

FIG. 28

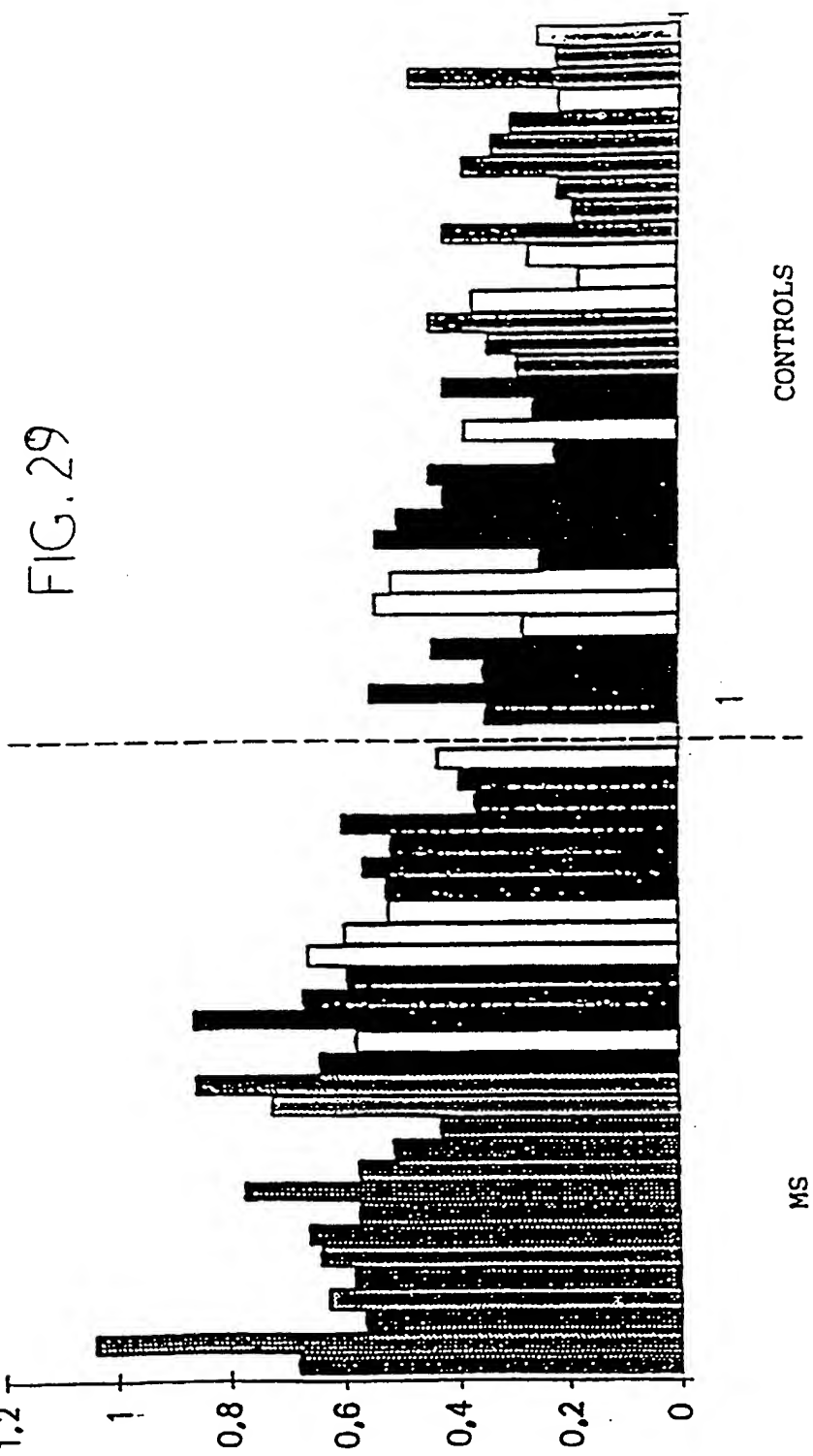
GATGCCTTTCTGCATCCCTGTACGTCTGACTCTCAATTCTTGCCTTGAAAG  
ATCCTTGAACCCAACGTCTCAACTCACCTGGACTGTTTACCCCAAGGGTTCAGGGA  
TAGCCCCATCTATTGGCCAGGCATTAGCCCAAGATGCCTTTGCATCCCTGTACGTG  
ACTCTCAATTCTTGCCTTGCCCTTGAAAGATGCTTGAACCCAACGTCTCAACT  
CACCTGGACTGTTTACGCCAAGGGTTCAGGGATAGCCCCCATCTATTGGC  
CAGGCATTAGCCCAA

SEQ ID NO 40

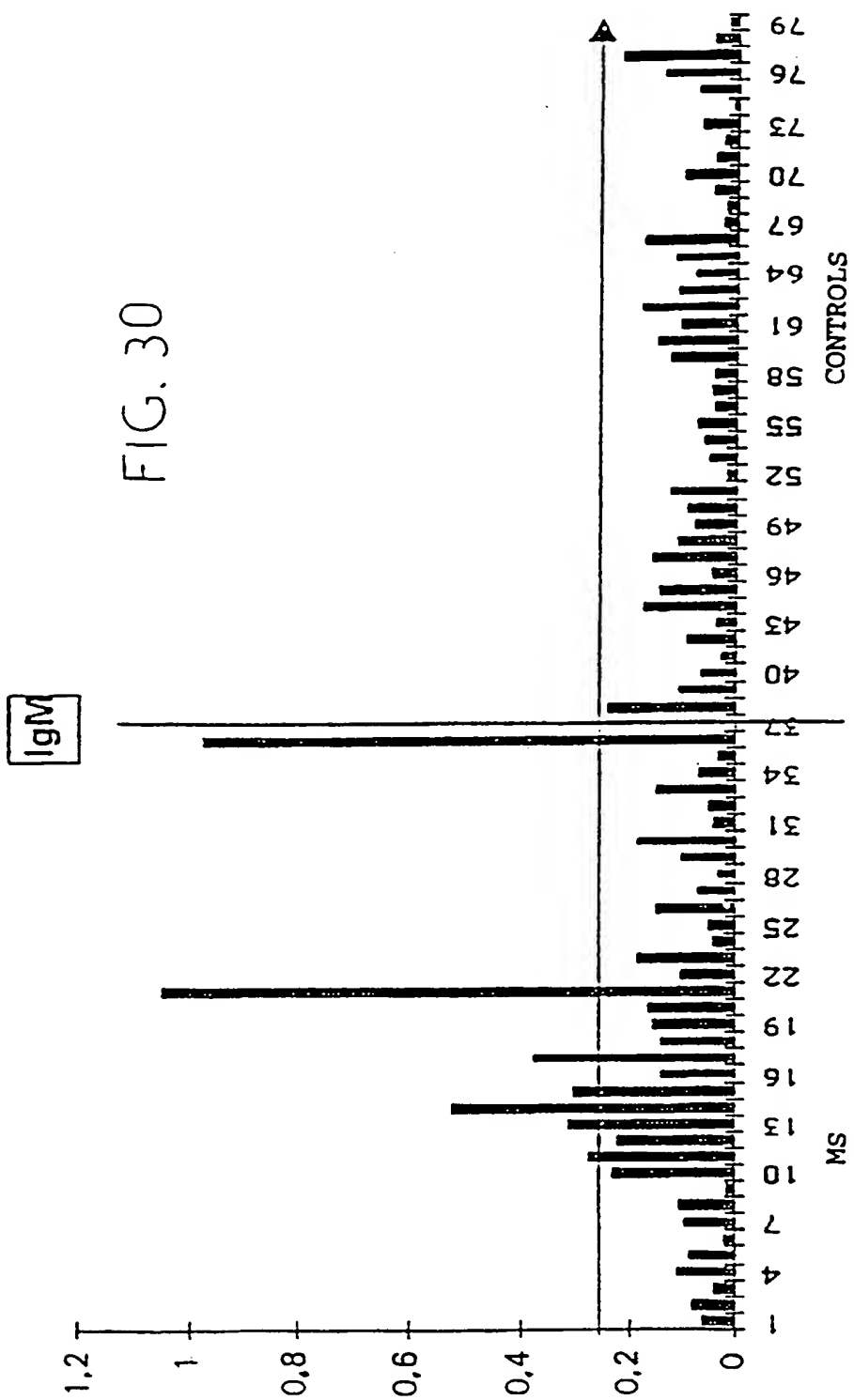
Asp-Ala-Phe-Phe-Cys-Ile-Pro-Val-Arg-Pro-Asp-Ser-Gln-Phe-  
Leu-Phe-Ala-Phe-Glu-Asp-Pro-Leu-Asn-Pro-Thr-Ser-Gln-Leu-  
Thr-Trp-Thr-Val-Leu-Pro-Gln-Gly-Phe-Arg-Asp-Ser-Pro-His-  
Leu-Phe-Gly-Gln-Ala-Leu-Ala-Gln

SEQ ID NO 39 (POL2B)

30/69

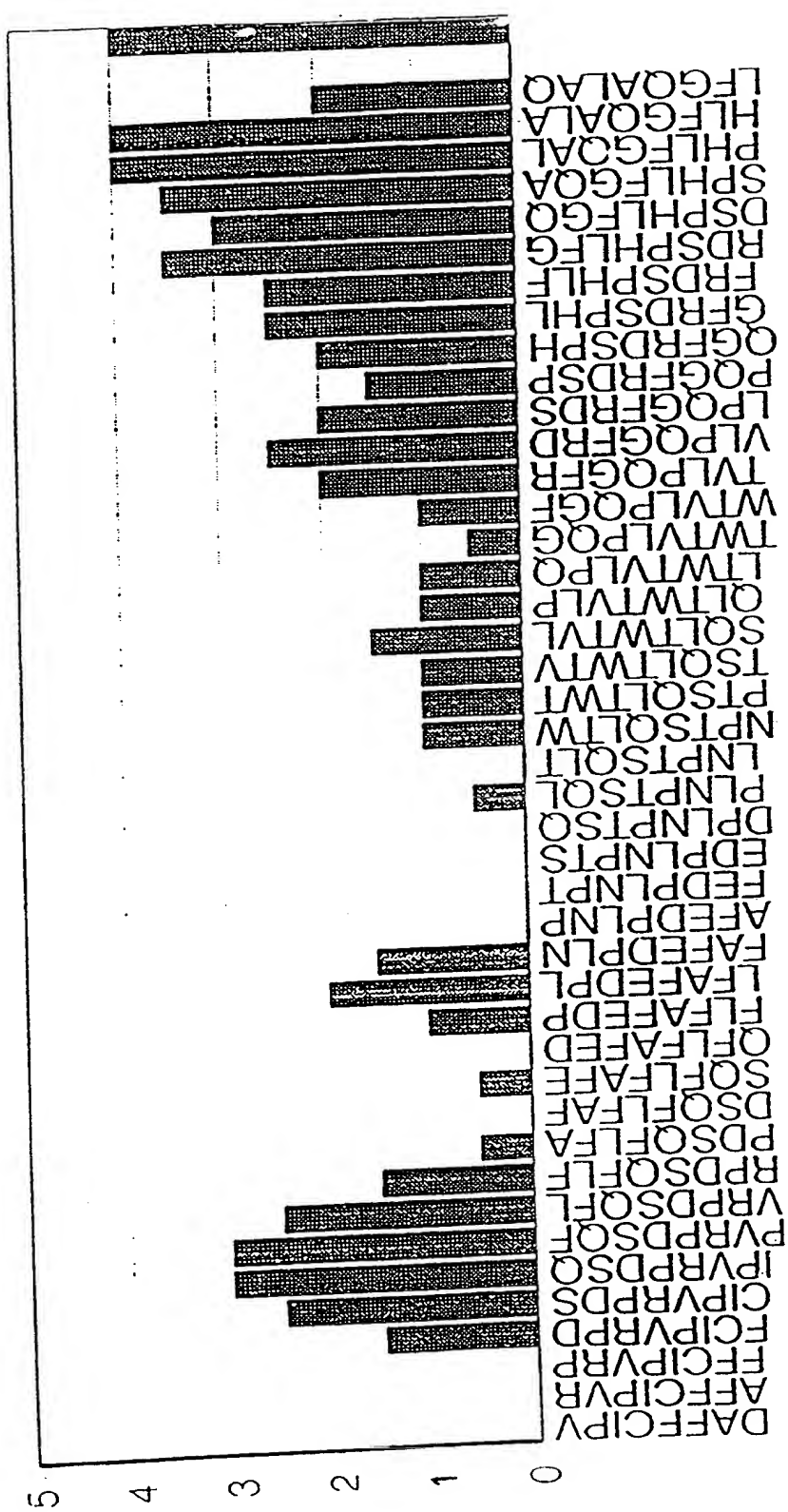


31/69



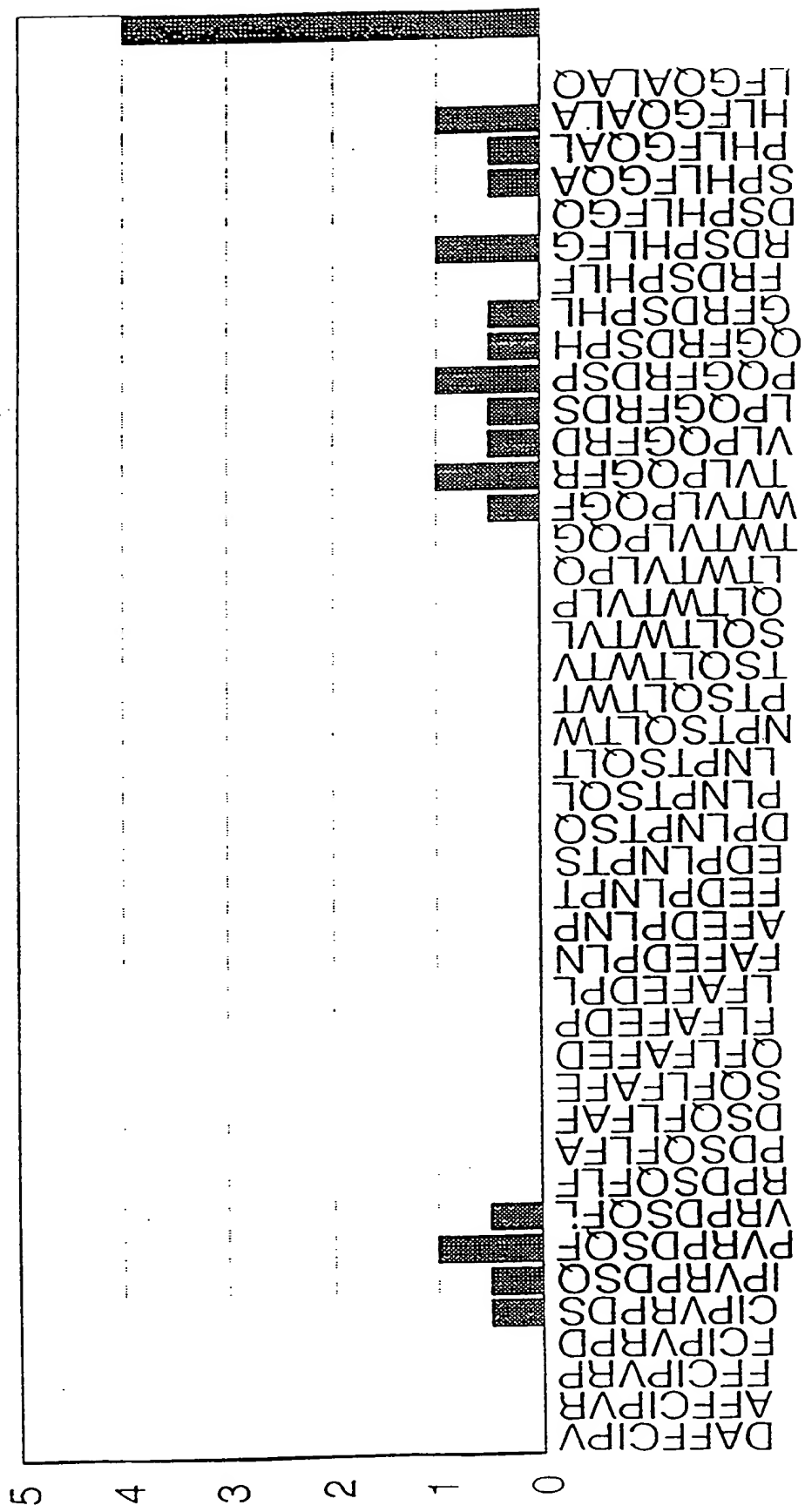
32/69

FIG. 31

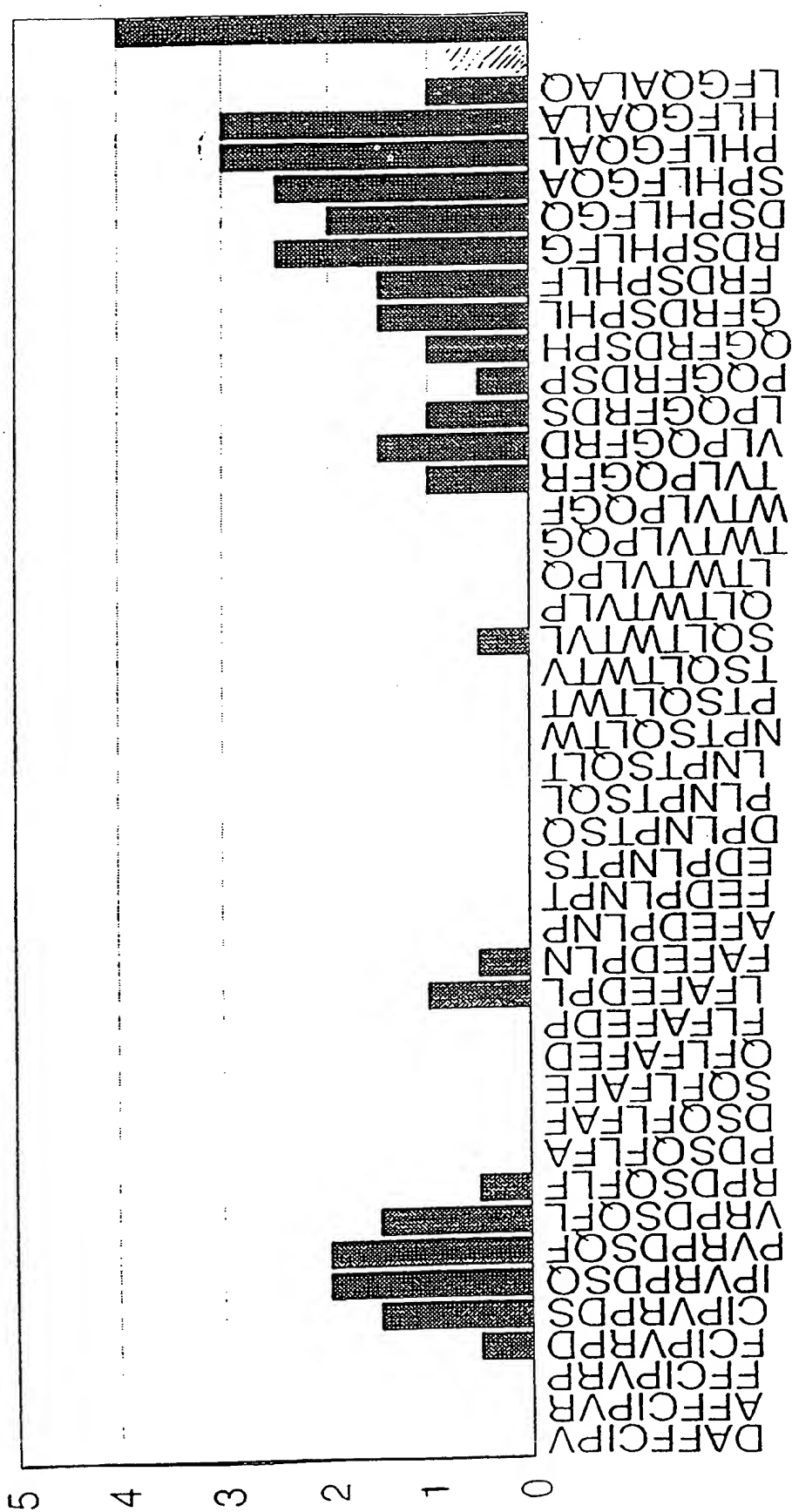


33/69

FIG. 32



34/69



*35/69*

## FIG. 34

Cys-Ile-Pro-Val-Arg-Pro-Asp-Ser-Gln-Phe-Leu SEQ ID NO 41

Val-Leu-Pro-Gln-Gly-Phe-Arg-Asp-Ser-Pro-His-Leu-Phe-Gly-  
Gln-Ala-Leu-Ala SEQ ID NO 42

Leu-Phe-Ala-Phe-Glu-Asp-Pro-Leu SEQ ID NO 43  
Phe-Ala-Phe-Glu-Asp-Pro-Leu-Asn SEQ ID NO 44

## FIG 35

36/69

10	20	30	40	50	
1234567890	1234567890	1234567890	1234567890	1234567890	100
CTTCCCCAAC	TAATAAGGAC	CCCCCITICA	ACCCAAACAG	TCCAAAAGGA	50
L P Q L	I R T	P L S	T Q T V	Q K D	
F P N . .	G P P F Q	P K Q	S K R T		
S P T N K D	P P F N	P N S	P K G		
CATAGACAAA	GGAGTAAACA	ATGAACCAAA	GAGTGCCAAT	ATTCCTGGT	100
I D K G V N N	E P K S A N	I P W L			
. T K E . T	M N Q R V P I	F P G			
H R Q R S K Q .	T K E C Q Y	S L V			
TATGCACCCCT	CCAAGCGGTG	GGAGAAGAAT	TOGGCCCAAGC	CAGAGTGGCAT	150
C T L Q A V G E E F	G P A R V H				
Y A P S K R W E K N	S A Q P E C M				
M H P P S G G R R I	R P S Q S A C				
GTACCTTTT	CCTCTCACCA	CTTGAAGCAA	ATTAATAG	ACNTAGGINA	200
V P F S L S H L K Q	I K I D X G X				
Y L F L S H T . S K L K .	T . V N				
T F F S L T L E A N .	N R X R X				
ATINICAGAT	AGCCCTGAATG	GYTATATTGA	TGTTTTACAA	GGATTAGGAC	250
X S D S P D G Y I D	V L Q G L G Q				
X Q I A L M X I L M	F Y K D . D				
I X R . P . W L Y .	C F T R I R T				
AATCCTTGA	TCTGACATGG	AGAGATATAA	TATTACTGCT	AAATCAGACG	300
S F D L T W R D I I	L L L N Q T				
N P L I . H G E I .	Y Y C . I R R				
I L . S D M E R Y N	I T A K S D A				
CTAACCTCAA	ATGAGAGAAG	TGCTGCCATA	ACTGGACCCC	GAGAGTTGG	350
L T S N E R S A A I	T G A R E F G				
. P Q M R E V L P .	L E P E S L A				
N L K . E K C C H N	W S P R V W				
CAATCTCTGG	TATCTCAGTC	AGGTCATGA	TAGGATGACA	ACGGAGGAAA	400
N L W Y L S Q V N D	R M T T E E R				
I S G I S V R S M I	G . Q R R K				
Q S L V S Q S G Q .	D D N G G K				
GAGAACGATT	CCCCACAGGG	CAGCAGGGAG	TTCGGAGTGT	ACCTCTCAT	450
E R F P T G Q Q A V	P S V A P H				
E N D S P Q G S R Q	F P V . L L I				
R T I P H R A A G S	S Q C S S S L				
TGGGACACAG	AATCAGAAC	TGGAGATTGG	TGCCGAGAC	ATTTA	495
W D T E S E H G D W	C R R H L				
G T Q N Q N M E I G	A A D I				
G H R I R T W R L V	P Q T F				

37/69

## FIG 36

10	20	30	40	50	
1234567890	1234567890	1234567890	1234567890	1234567890	
C T T C C C A A C	T A A T A A G G A C	C C O C C T T C A	A C C C A A C A G	T C C A A A A Q G A	50
L P Q L	I R T	P L S	T Q T V	Q K D	
C A T A G A C A A A	G G A G T A A A C A	A T G A A C C A A A	G A G T G C C A A T	A T T C C C I C G T	100
I D K	G V N N	E P K	S A N	I P W L	
T A T G C A C C C T	C C A A G O G G T G	G G A G A A G A A T	T C G G C C C A G C	C A G A G T C C A T	150
C T L	Q A V	G E E F	G P A	R V H	
G T A C C T T T T T	C I C T C T C A C A	C T T G A A G C A A	A T T A A A A T A G	A C C T A G G T A A	200
V P F S	L S H	L K Q	I K I D	L G K	
A T I C T C A G A T	A G C C C T G A T G	G Y T A T A T T G A	T G T T T T A C A A	G G A T T A G G A C	250
F S D	S P D G	Y I D	V L Q	G L G Q	
A A T C C T T T G A	T C T G A C A T G G	A C A G A T A T A A	T A T T A C T G C T	A A A T C A G A C G	300
S F D	L T W	R D I I	L L L	N Q T	
C T A A C C T C A A	A T G A G A G A A G	T G C T G C C A T A	A C T G G A C C C	G A G A G T T T G G	350
L T S N	E R S	A A I	T G A R	E F G	
C A A T C T C I G G	T A T C T C A G I C	A G G T C A A T G A	T A G G A T G A C A	A C G G A G G A A A	400
N L W	Y L S Q	V N D	R M T	T E E R	
G A G A A C G A T T	O O C C A C A G E G G	C A G C A G G C A G	T T C C A G I G T	A G C T C C T C A T	450
E R F	P T G	Q Q A V	P S V	A P H	
T G G G A C A C A G	A A T C A G A A C A	T G G G A G T T G G	T G C O G C A G A C	A T T T A C A A C T	500
W D T E	S E H	G D W	C R R H	L Q L	
T G G T G C T A N	A A G G A C T I N A G	G A A A C T I A G G	A A G A C T I A N G A	A T T A T C A A N	550
A C X	K D X G	K L G	R L X	I I Q X	
G A T G T C C A C T	A N N A C A C A G G	G G A A A G G A A G	A A A A T C C T A C	T G C T T T T C T G	600
C P L	X H R	G K E E	N P T	A F L	
G A G A G A C T A A	G G G A G G C A T T	G A G G A A G C A T	A C C A G G C A A G	T G G A C A T T G G	650
E R L R	E A L	R K H	T R Q V	D I G	
A G G C T C T G G A	A A A G G G A A A A	G T T G G G C A A A	T T A T A T G C T	A A T A C G G C T	700
G S G	K G K S	W A N	Y M P	N R A C	
G C T T O C A G I G	C A G I C T A C A A	G G A O G C T T T A	G A A A G A T T G	T C C A A G T A G A	750
F Q C	S L Q	G R F R	K D C	P S R	
A A T A A G C C G C	C C C T C G T C C A	T G C C C C T T A T	G T C A A G G G A A	T C A C T G G A A G	800
N K P P	L V H	A P Y	V K G I	T G R	
G C C T A C T G C C	O C A G G G G A C G	A A G G T C C T C T	G A G T C A G A A G	C C A C T A A C C T	850
P T A	P G D E	G P L	S Q K	P L T	
GA					852

38/69

FIG 37

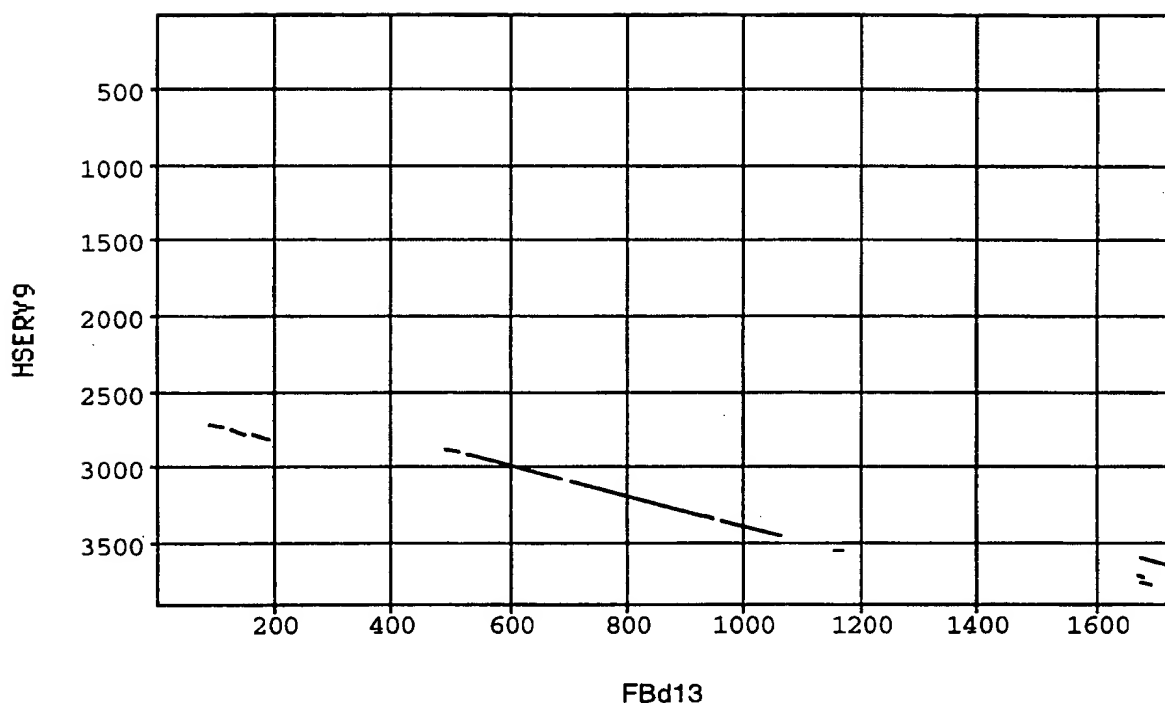


FIG 38  
G

39/69

10	20	30	40	50	
<u>1234567890</u>	<u>1234567890</u>	<u>1234567890</u>	<u>1234567890</u>	<u>1234567890</u>	
AAGGAAACTC	AGAAAGCCAA	TACCCATTAA	GTAAGATGGA	CACCAAGC	50
K E T Q	K A N	T H L	V R W T	P E A	
R K L	R K P I	P I .	. D G	H Q K Q	
G N S	E S Q	Y P F S	K M D	T R S	
AGAAGCAGCT	TICCAGGCC	AAAGAAAATC	CCTAACCCAA	GCCCCAGTGT	100
E A A	F Q A L	K K S	L T Q	A P V L	
K Q L	S R P	. R N P	. P K	P Q C	
R S S F	P G P	K E I	P N P S	P S V	
TAAGCTTGC	AAACGGGCAA	GACTTTCTT	TATAATGTCAC	AGAAAAACAG	150
S L P	T G Q	D F S L	Y V T	E K Q	
. A C Q	R G K	T F L	Y M S Q	K N R	
K L A	N G A R	L F F	I C H	R K T G	
GAATAGCTCT	AGGAGTCCTT	ACACAGGTCC	AAGGGACAAG	CTTGCACCT	200
E . L .	E S L	H R S	K G Q A	C N L	
N S S	R S P Y	T G P	R D K	L A T C	
I A L	G V L	T Q V Q	G T S	L Q P	
GTGGCATACC	TGAGTAAGGA	AACTGATGTA	NTGGCAAAGG	GTGGCCTCA	250
W H T	. V R K	L M X	W Q R	V G L I	
G I P E	. G N .	C X	G K G	L A S	
V A Y L	S K E	T D V	X A K G	W P H	
T T G T T A C A G	G T A G G G C A G C	A G T A G C A G T C	T T A G T T T C T G	A A A C A G T T A A	300
V Y R	. G S	S S S L	S F	. N S .	
L F T G	R A A	V A V	L V S E	T V K	
C L Q	V G Q Q	. Q S	. F L	K Q L K	
A A T A A T A C A G	G G A A G A G A T C	T T A C T G T G T G	G A C A T C T C A T	G A T G T G A A O G	350
N N T G	K R S	Y C V	D I S .	C E R	
I I Q	G R D L	T V W	T S H	D V N G	
. Y R	E E I	L L C G	H L M M	. T	
G C A T A C T C A C	T G C T A A A G A G	G A C T T G T G C	T G T C A G A C A A	C C A T T A C T T	400
H T H C	. R G L V A	V R Q	P F T	; I L T A K E	
I L T	D L W L	S D N	H L L		
A Y S L	L K R	T C G	C Q T T	I Y L .	
A A A T A G C A G G	T T C T A T T A C T	T G A A G T G C C A	G T G C T G O G A C	T G C A C A T T T G	450
I A G S I T	. S A S A A T	A H L			
K . Q V	L L L	E V P	V L R L	H I C	
N S R	F Y Y L	K C Q	C C D	C T F V	
T G C A A C T C T T	A A C C C A G C C A	C A T T C T T C C	A G A C A A T G A A	G A A A G A T A G	500
C N S .	P S H	I S S	R Q .	R K D R	
A T L	N P A T	F L P	D N E	E K I E	
Q L L	T Q P	H F F Q	T M K	K R .	

FIG38

b

40/69

10	20	30	40	50	
<u>1234567890</u>	<u>1234567890</u>	<u>1234567890</u>	<u>1234567890</u>	<u>1234567890</u>	
AACATAACTG	TCAACAAGTA	ATTGCTAAA	CCTATGCTGC	TOGAGGGGAC	550
T . L	S T S N	C S N	L C C	S R G P	
H N C	Q Q V	I A Q T	Y A A	R G D	
N I T V	N K .	L L K	P M L L	E G T	
CTTCTAGAGG	TTCCCTTGAC	TGATCCGAC	CTCAACTTGT	ATACTGATGG	600
S R G	S L D	. S R P	Q L V	Y . W	
L L E V	P L T	D P D	L N L Y	T D G	
F . R F P	. L I P T	S T C	I L M E		
AAGITCCTTG	GCAGAAAAAG	GACTTTGAAA	ACCGGGGTAT	GCAGTGTATCA	650
K F L G	R K R	T L K	S G V C	S D Q	
S S L	A E K G	L . K	A G Y	A V I S	
V P W	Q K K	D F E K	R G M	Q . S	
GIGATAATGG	AATACTTGAA	AGTAATGCC	TCACTCCAGG	AACTAGTGCT	700
. . W N T	. K . S P	H S R N	. C S		
D N G	I L E	S N R L	T P G	T S A	
V I M E	Y L K	V I A	S L Q E	L V L	
CACCTGGCAG	AACTAATAGC	CCTCACTTGG	GCACTAGAAT	TAGGAGAAGG	750
P G R	T N S	P H L G	T R I	R R R	
H L A E	L I A	L T W	A L E L	G E G	
T W Q	N . . P	S L G	H . N .	E K E	
AAAAAGGGTA	AATATATATT	CAGACTCTAA	GTATGCTTAC	CTAGTCCTCC	800
K K G K	Y I F	R L .	V C L P	S P P	
K R V	N I Y S	D S K	Y A Y	L V L H	
K G .	I Y I	Q T L S	M L T .	S S	
ATGCCCATGC	AGCAATATGG	AGAGAGAGGG	AATTCCTAAC	TTCCTGAGGGA	850
C P C	S N M E	R E G	I P N	F . G N	
A H A	A I W	R E R E	F L T	S E G	
M P M Q	Q Y G	E R G	N S .	L L R E	
ACACCTATCA	ACCATCAGGG	AAGCCATTAG	GAGATTATTA	TTGGCTGTAC	900
T Y Q	P S G	K P L G	D Y Y	W L Y	
T P I N	H Q G	S H .	E I I I	G C T	
H L S	T I R E	A I R	R L L	L A V Q	
AGAAACCTAA	AGAGGTGGCA	GTCTTACACT	GCCAGGGTCA	TCAGGAAGAA	950
R N L K	R W Q	S Y T	A R V I	R K K	
E T .	R G G S	L T L	P G S	S G R R	
K P K	E V A	V L H C	Q G H	Q E E	
GAGGAAAGGG	AAATAGAAGG	CAATCGCAA	CGGGATATTG	AAGCAAAAAA	1000
R K G	K . K A	I A K	R I L	K Q K K	
G K G	N R R	Q S P S	G Y .	S K K	
E E R E	I E G	N R Q	A D I E	A K K	

*4/16/98*

**FIG 38**  
C

10	20	30	40	50	
1234567890	1234567890	1234567890	1234567890	1234567890	1234567890
AGCCCCAAGG CAGGACTCTC CATTAGAAAT GCTTATAGAA GGACCCCTAG					1050
P Q G R T L H . K C L . K D P .					
S R K A G L S I R N A Y R R T P S					
A A R Q D S P L E M L I E G P L V					
TATGGGGTAA TCCCCCTCTGG GAAACCAAGC CCCAGTACTC ACCAGGAAAA					1100
Y G V I P S G K P S P S T Q Q E K					
M G . S P L G N Q A P V L S R K N					
W G N P L W E T K P Q Y S A G K					
ATAGAACATAGG AAACCTCACA AGGACATACT TTCCCTCCCT CCAGATGGCT					1150
. N R K P H K D I L S S P P D G .					
R I G N L T R T Y F P P L Q M A					
I E . E T S Q G H T F L P S R W L					
AGCCACTGAG GAAGGAA					1167
P L R K E					
S H . G R					
A T E E G					

*42/69*

FIG 39

10	20	30	40	50	
1234567890	1234567890	1234567890	1234567890	1234567890	
AACTTGCGTG	CTAGAACGAC	TAAGGAAAAC	TAGGAAGACT	ATGAATTATT	50
N L R A	R R T	K E N .	E D Y	E L F	
T C V	L E G L	R K T	R K T	M N Y S	
L A C .	K D .	G K L	G R L .	I I	
<b>C</b>	CAATGATGTC	CACTATAACA	CAGGGGAAAG	GAAGAAAATC	100
N D V	H Y N T	G E R K K I	L L P F		
M M S	T I T	Q G K G	R K S	Y C L	
Q .	C P L .	H R G K	E E N P	T A F	
TCTGGAGAGA	CTAAGGGAGG	CATTGAGGAA	GCATACCAAG	CAAGTGGACA	150
W R D .	G R H .	G S I P G	K W T		
S G E T	K G G I E E	A Y Q A	S G H		
L E R	L R E A	L R K	H T R	Q V D I	
TTGGAGGCTC	TGGAAAAGGG	AAAAGTGGG	CAAATTGAAT	GCCTAATAGG	200
L E A L	E K G K V G	Q I E C	L I G		
W R L	W K R E	K L G K L N A . . G			
G G S	G K G K S W A	N . M P N R			
GCTTGCTTCC	AGTGCAGTCT	ACAAGGACGC	TTTAGAAAAG	ATITGCCAAG	250
L A S	S A V Y	K D A L E K	I V Q V		
L L P	V Q S T R T L .	K R L S K			
A C F Q	C S L Q G R	F R K D C P S			
TAGAAATAAG	CGCGCCCTCG	TCCATGCCCC	TTATGTCAAG	GGAATCACTIG	300
E I S	R P S S M P L	M S R E S L			
. K .	A A P R P C P	L C Q G N H W			
R N K	P P L V H A P	Y V K G I T G			
GAAGGCCATAC	TGCCCCACGG	GACGAAGGTC	CTCTGAGTCA	GAAGCCACTA	350
E G L L	P Q G T K V L .	V R S H .			
K A Y C P R G	R R S S E S	E A T N			
R P T A P G D E G P	L S Q K P L				
ACCTGATGAT	CCAGCAGCAG	GACTGAGGGT	GGGGGGGCA	AGTGCAGGC	400
P D D	P A A G L R V	P G A S A S P			
L M I	Q Q Q D .	G C P G Q V P A			
T . .	S S S R T E G	A R G K C Q P			
CATGCCATCA	CCCTCAGAGC	CCGGGTATG	TTTGACCATT	GAGGCCAGG	450
C H H	P Q S P G Y V .	P L R A R			
H A I T	L R A P G M	F D H . E P G			
M P S	P S E P R V C	L T I E S Q E			
AAGTTAACIG	TCTCTGGAC	ACTGGGCCAG	CTTCTTCAGT	CTTACTTTCC	500
K L T V	S W T L A Q	P S Q S Y F P			
S . L	S P G H W R S	L L S L T F L			
V N C	L L D T G A A	F S V L L S			

43/59

FIG 39  
b

10	20	30	40	50	
1234567890	1234567890	1234567890	1234567890	1234567890	
TGTCCCCAGAC	AATTGTCCTC	CAGATCTGTC	ACTATCOGAG	GGGTCCCTAAG	550
V P D N C P P	D L S L S E G S . D				
S Q T I V L Q I C H Y P R G P K					
C P R Q L S S R S V T I R G V L R					
ACAGGCCAGTC	ACTACATACT	TCTCTCAGCC	ACTAAGTGT	GACTGGGGAA	600
S Q S L H T S L S H . V V T G E					
T A S H Y I L L S A T K L . L G N					
Q P V T T Y F S Q P L S C D W G T					
CTTTACTCTT	TTCACATGCT	TTTCTAATT	TGCCTGAAAG	CCCCACTCC	650
L Y S F H M L F . L C L K A P L P					
F T L F T C F S N Y A . K P H S L					
L L F S H A F L I M P E S P T P					
TIGTTAGGGA	GAGACATTTT	AGCAAAAGCA	GGGGCCATTA	TACACCTGAA	700
C . G E T F . Q K Q G P L Y T . T					
V R E R H F S K S R G H Y T P E					
L L G R D I L A K A G A I I H L N					
CATAGAAAAA	GGAATACCCA	TTTGCTGTCC	OCTGCCTGAG	GAAGGAATT	750
. E K E Y P F A V P C L R K E L					
H R K R N T H L L S P A . G R N .					
I G K G I P I C C P L L E E G I N					
ATCCTGAAGT	CTGGGCAATA	GAAGGACAAT	ATGGACAAGC	AAAGAATGCC	800
I L K S G Q . K D N M D K Q R M P					
S . S L G N R R T I W T S K E C P					
P E V W A I E G Q Y G Q A K N A					
CGTCTGTTC	AAGTTAAACT	AAAGGATCT	CCCTCTTTC	CCTACCAAAG	850
V L F K L N . R I L P P F P T K G					
S C S S . T K G F C L L S L P K					
R P V Q V K L K D S A S F P Y Q R					
GAAGTACCCP	CTTAGACCCG	AGGCCCTACA	AGGACTCAAA	AGATTGTTAA	900
S T L L D P R P Y K D S K D C .					
E V P S . T R G P T R T Q K I V K					
K Y P L R P E A L Q G L K R L L R					
GGACCTAAAA	GCCCAAGGCC	TAGTAAAACC	ATGCAGTAGC	CCCTIGCAATA	950
G P K S P R P S K T M Q . P L Q Y					
D L K A Q G L V K P C S S P C N T					
T . K P K A . . N H A V A P A I					
CTCCAATT	AGGAGTAAGG	AAACCCAACG	GACAGTGGAG	GTTAGTGCAA	1000
S N F R S K E T Q R T V E V S A R					
P I L G V R K P N G Q W R L V Q					
L Q F . E . G N P T D S G G . C K					

44169

FIG 39  
C

10	20	30	40	50	
1234567890	1234567890	1234567890	1234567890	1234567890	
GATCTCACCA TTATTAATGA GGCIGTTTT CCTCTATAACC CAGCTGTATC					1050
S Q D Y . . G C F S S I P S C I					
D L R I I N E A V F P L Y P A V S					
I S G L L M R L F F L Y T Q L Y L					
TAGCCCCTTAT ACTCTGCTTT CCTTAATACC AGAGGAACCA GAGTAGTTA					1100
. P L Y S A F P N T R G S R V V Y					
S P Y T L L S L I P E E A E . F T					
A L I L C F P . Y Q R K Q S S L					
CAGTCCTCGGA CCTTAAGGAT GCCTCTTTCT GCATCCCTGT ACATCCTGAT					1150
S P G P . G C L F L H P C T S . F					
V L D L K D A S F C I P V H P D					
Q S W T L R M P L S A S L Y I L I					
TCTCAATTCT TGTTTGCTT TGAAGATCT TTGAACCCAA TGTCICAATT					1200
S I L V C L . R S F E P N V S I					
S Q F L F V F E D P L N P M S Q F					
L N S C L S L K I L . T Q C L N S					
CACCTGGACT GTTTACCCC AGGGGTTCCG GGATAGCCCC CATCTATTIG					1250
H L D C F T P G V P G . P P S I W					
T W T V L P Q G F R D S P H L F G					
P G L F Y P R G S G I A P I Y L					
GCCAGGCATT AGCCCAAGAC TTGACCAAT TCTCATACCT GGACATCTTG					1300
P G I S P R L E P I L I P G H L V					
Q A L A Q D L S Q F S Y L D I L					
A R H . P K T . A N S H T W T S C					
TOCTTGGTA TGGGATGATT TAATTTAGC CACCGCTCA GAAACCTTGT					1350
L R Y G M I . F . P P V Q K P C					
S F G M G . F N F S H P F R N L V					
P S V W D D L I L A T R S E T L C					
GCCATCAAGC CAACCAAGCG TICITAAATT TOCTCACTCC GTGIGGCTAC					1400
A I K P P K R S . I S S L R V A T					
P S S H P S V L K F P H S V W L Q					
H Q A T Q A F L N F L T P C G Y					
AAGGTTCCA AACCAAAGGC TCAGCTCTGC TCACAGCAGG TAAATACCT					1450
R F P N Q R L S S A H S R L N T .					
G F Q T K G S A L L T A G . I L					
K V S K P K A Q L C S Q Q V K Y L					
AGGGTTAAAA TTATCAAAG GCACCAGGGC CCTCTGTGAG GAATGTATCC					1500
G . N Y P K A P G P S V R N V S					
R V K I I Q R H Q G P L . G M Y P					
G L K L S K G T R A L C E E C I Q					

45/69

FIG 39  
d

10	20	30	40	50
1234567890	1234567890	1234567890	1234567890	1234567890
AACCTGTACT	GGCTTATCIT	CATCCAAAAA	CCCTAAAGCA	ACTAAGAAGG
N L Y W	L I F	I P K	P . S N	. E G
T C T	G L S S	S Q N	P K A	T K K V
P V L	A Y L	H P K T	L K Q	L R R

1550

TCCTTGGCAT AACAGGTTTC TGCGGAA 1577  
P W H N R F L P  
L G I T G F C R  
S L A . Q V S A E

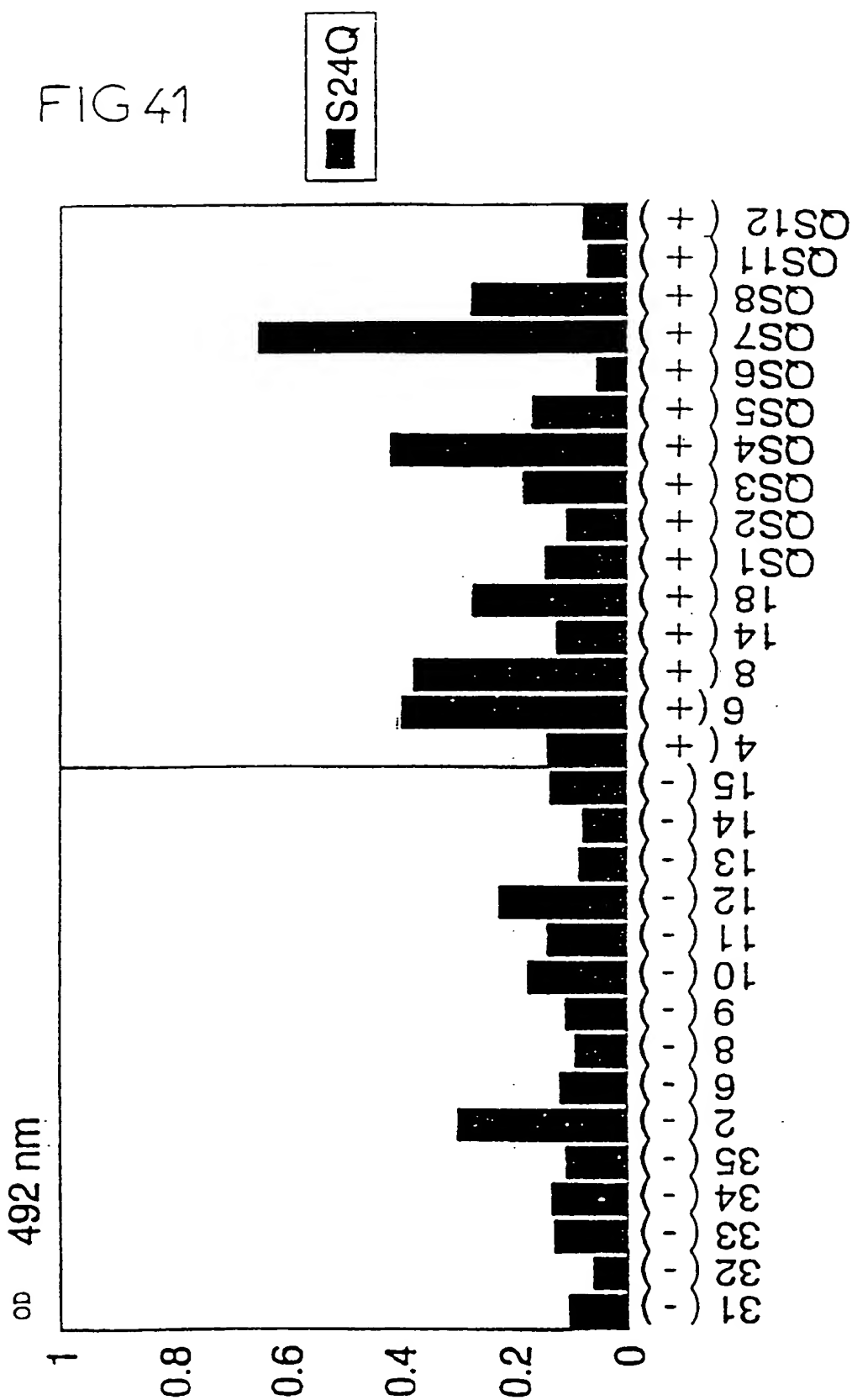
46/69

## FIG 40

10	20	30	40	50	
<u>1234567890</u>	<u>1234567890</u>	<u>1234567890</u>	<u>1234567890</u>	<u>1234567890</u>	
TCCAGCAGCA	GGACTGAGGG	TGCCCGGGGC	AAGTGCCAGC	CCATGCCATC	50
S S S R	T E G	A R G	K C Q P	M P S	
ACCCTCAGAG	CCCCGGGTAT	GTTTGACCAT	TGAGAGCCAG	GAAGTTAACT	100
P S E	P R V C	L T I	E S Q	E V N C	
GTCCTCTGGA	CACTGGCGCA	GCCTTCTCAG	TCTTACTTTC	CTGTCCCCAGA	150
L L D	T G A	A F S V	L L S	C P R	
CAATTGTCCT	CCAGATCTGT	CACTATCGA	GGGGCTCTAA	GACAGCCAGT	200
Q L S S	R S V	T I R	G V L R	Q P V	
CACTACATAC	TCTCTCTCAGC	CACTAAGTTG	TGACTGGGA	ACTTTACTCT	250
T T Y	F S Q P	L S C	D W G	T L L F	
TTTCACATGC	TTTCTTAATT	ATGCTGAAA	GCCCCACTCC	CTTGTAGGG	300
S H A	F L I	M P E S	P T P	L L G	
AGAGACATTT	TAGAAAAGC	AGGGGCCATT	ATACACCTGA	ACATAGGAAA	350
R D I L	A K A	G A I	I H L N	I G K	
AGGAATACCC	ATTTGCTGTC	CCCTGCTTGA	GGAGGAATT	AATCTGAAG	400
G I P	I C C P	L L E	E G I	N P E V	
TCTGGCAAT	AGAAGGACAA	TATGGACAAG	CAAAGAATGC	CGTCTGTT	450
W A I	E G Q	Y G Q A	K N A	R P V	
CAAGTTAAC	TAAAGGATTIC	TGCCTCCITT	CCCTACCAAA	GGAGTACCC	500
Q V K L	K D S	A S F	P Y Q R	K Y P	
TCTTAGACCC	GAGGCCCTAC	AAGGACTCAA	AAGATTGTTA	AGGACCT	547
L R P	E A L Q	G L K	R L L	R T	

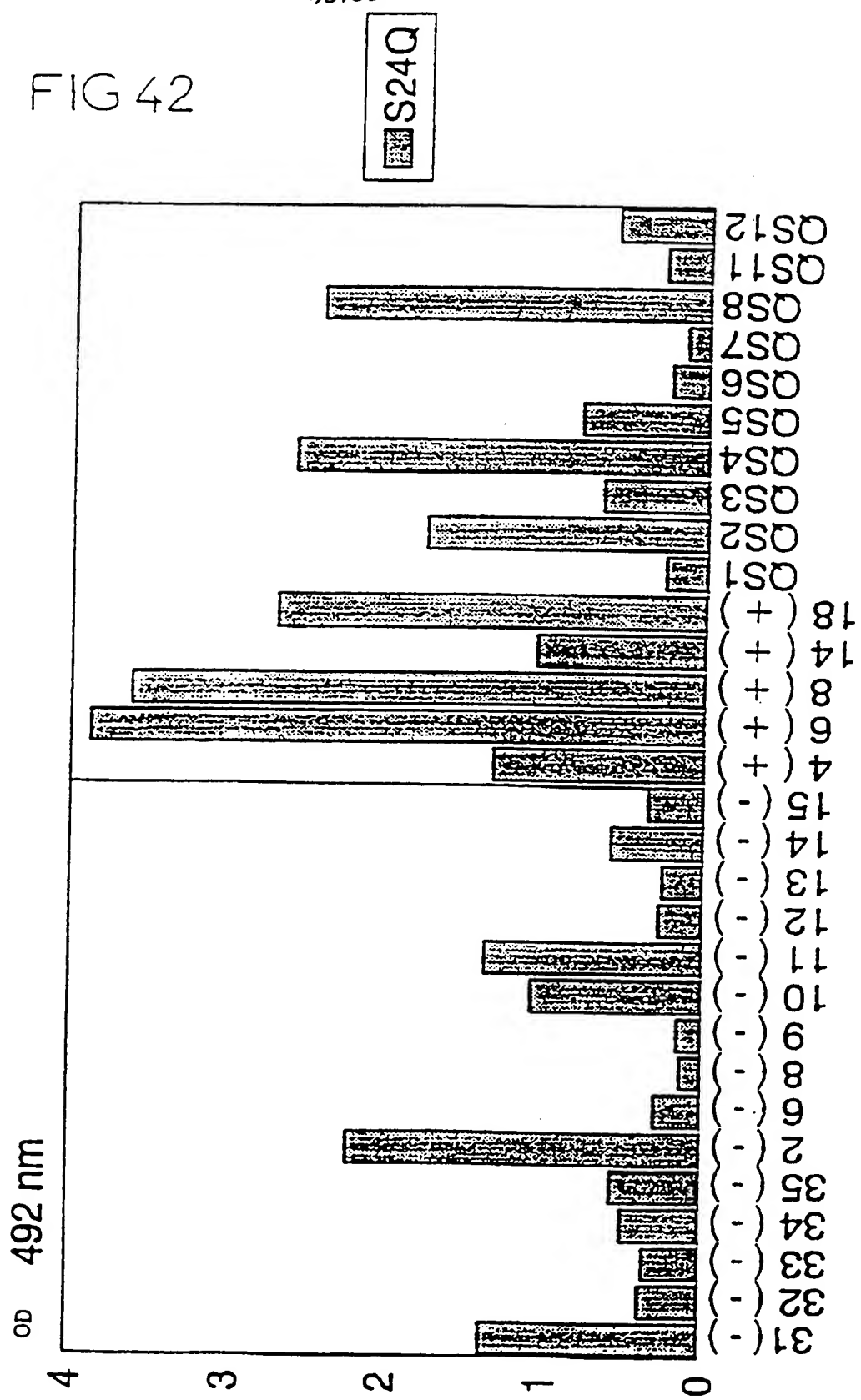
47/69

FIG 41



48169

FIG 42



49/69

FIG 43

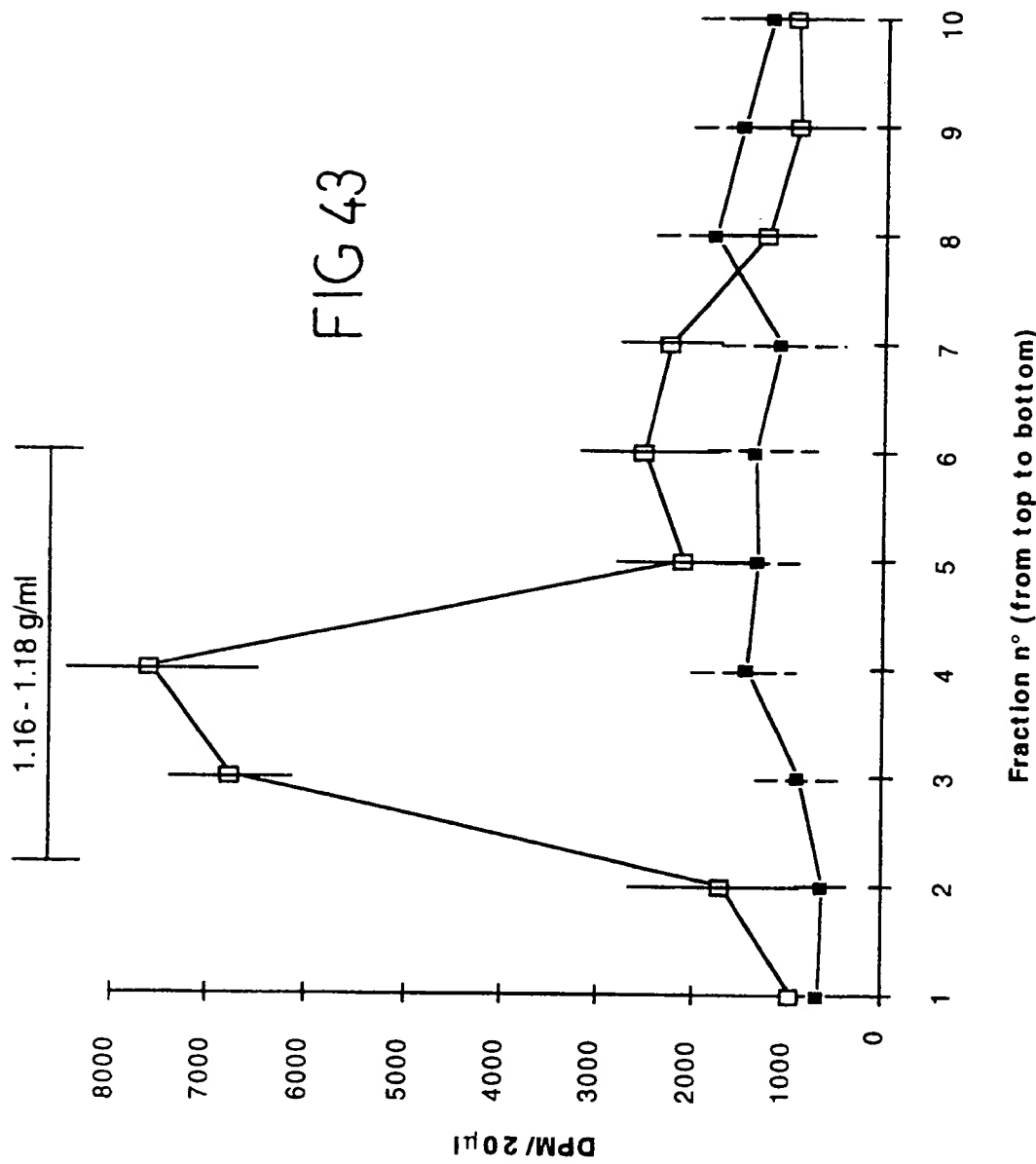
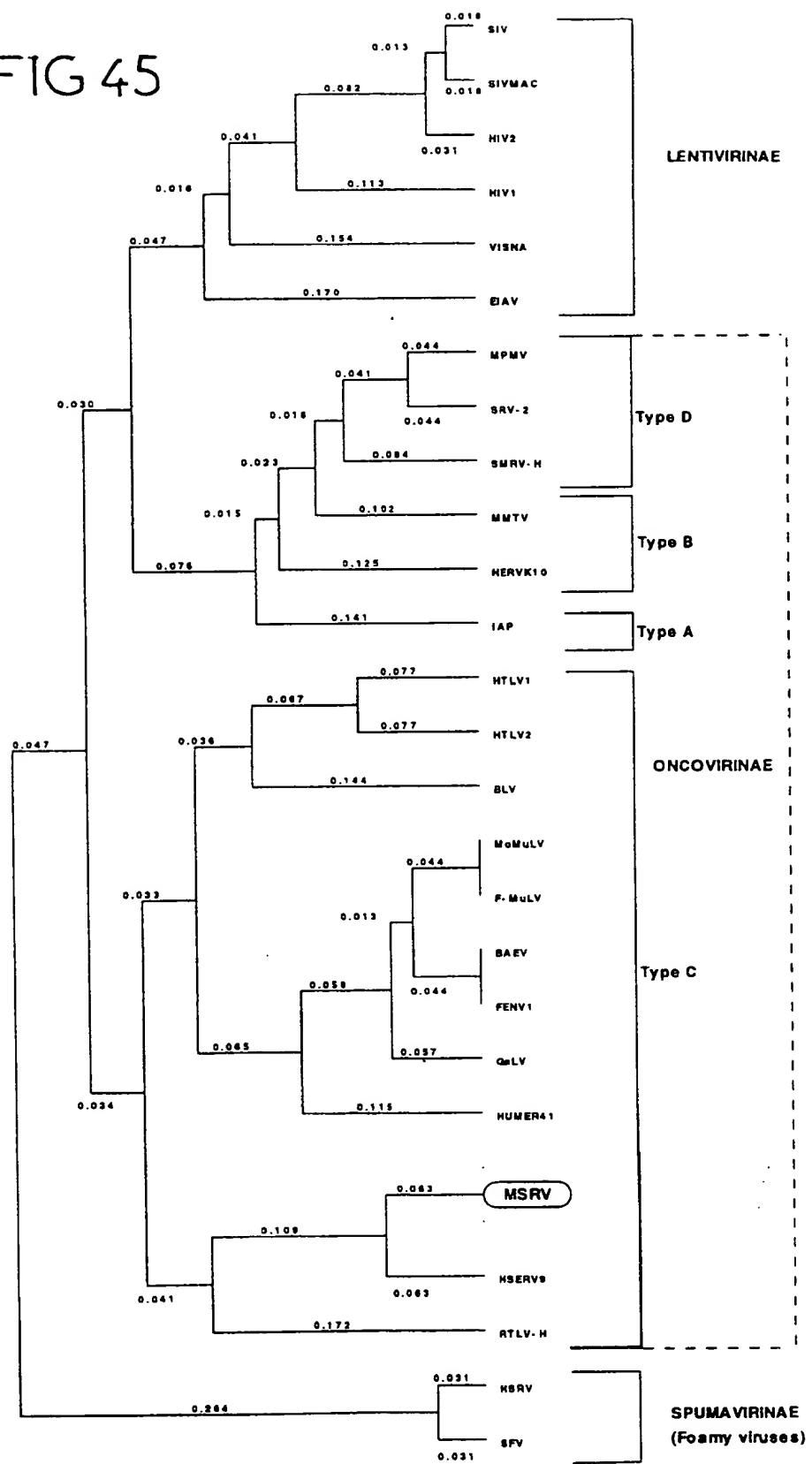


FIG 44

51/69

FIG 45



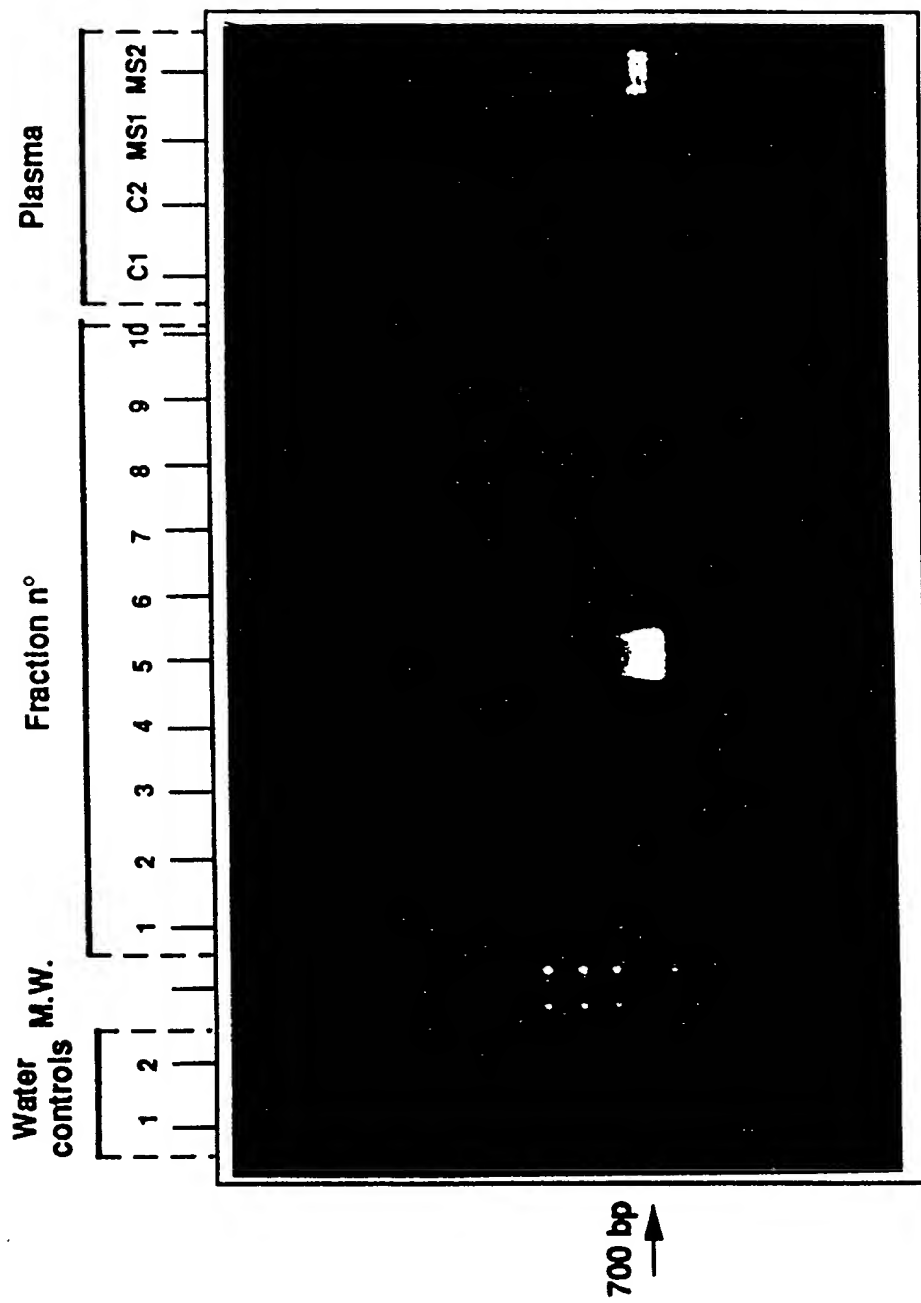
52/59

FIG 46

PROTEASE	TCCAGGACCA CGACTGGGGG TCCCGGGC AAGTCCACC CCTAGGATC G A R G K C Q P M P S	50	
	GCGCTAAGG CGCGCGAT GTTGTGACGT TGAGACCGAG GAAGTTAACT P S E P R V C L T I E S Q E V N C	100	
	GTCCTCGG CACTGGGGCA GCGTCTAG TCTTACTTC CTGTCGCCAGA L L D T G A A F S V L L S C P R	150	
	CAATTCTCT CCAGATGCT CACTATCGA CGGTCTAG GACGCCAGT Q L S S R S V T I R G V L G Q P V	200	
	CACTAGATC TTCTCTAGC CACTAGTTG TGACTGGGA ACTTTACTCT T T Y F S Q P L S C D W G T L L F	250	
	TTTCAGATC TTTCATATT ATGGCTGAA CGCCACTAC CTGCTAGG S H A F L I M P E S P T P L L G	300	
	AGAGACATTC TAGGAAAGC AGGGCGATC ATACACCTG ACAGTGAAA R D I L A K A G A I I H L N I G K	350	
	AGGAATCCC ATTTGCGTC CGCTGCTGAA GGAGGAATT AATCTGAG G I P I C C P L L E E G I N P E V	400	
	TCTGGCAAT AGAAGGCAA TTGGACAG CAAAGAACCC CGCTCTGTT W A I E G Q Y G Q A K N A R P V	450	
	CAAGTAAAC TAAGGATTC TGCTCTTCT CGTACCAA CGAAGTACCC Q V K L K D S A S F P Y Q R K Y P	500	R
	TCTTAGGCC GAGGCGTAC ARGANCICA AAGATGTT AAGGCGTAA L R P E A L Q G X Q K I V K D L K	550	E
	AAGCGCGAG CGCTAGTAAA CGTGCAGA CGCTCTGAA TACCTCAATT A O G L V K P C S S P C N T P I	600	V
	TTAGGGTA GGAAACCAA CGCACAGGG AGGTGATCC AAGCTCTAG L G V R K P N G Q W R L V Q D L R region A	650	E
	GATTTAAAT GAGCTGTTT TCTCTCTATA CGCGCTGTA TUGACCTT T I N E A V F P L Y P A V S S P Y	700	R
	ATACCTGCT TTCTCTATA CGAGAGGG CAGAGTGGTT TACAGTCTTG T L L S L I P E E A E W F T V L	750	S
	GACCTTAAAGG ATGCTTTTCT CGCTCATCT GTAGCTGGC ACTCTCAATT D L K D A F F C I P V R P D S Q F	800	C
	CTTGTTCG TTGAGAGTC CTGAGACCC AACGCTCTAA CTACCTGGA L F A F E D P L N P T S Q L T W T	850	R
	CCTTTTAC CGAACGGTC AGGGATACCC CGCTCTTATG TGCCAGGCA V L P Q G F R D S P H L F G Q A	900	A
	TTAGGCCAG ACTTGAGTC ATTCCTCTAC CGGACACAC TGCGCTCCA L A Q D L S Q F S Y L D T L V L Q	950	&
	GTAAGGCGAT GATTCTCTT TGTGCGGT TTGAGAACG TGCGCTCCT Y V D D L L L V A R S E T L C H Q	1000	N
	AGCGACCCA AGAACCTCTA ATCTTCCCA CTACCTGGG CTACAGGTT A T Q E L L T F L T T C G Y K V	1050	a
	TOCAACCAA AGGCTCGGT CTGCTCAGG GAGATGAGT ACTTGGCT S K P K A R L C S Q E T R Y L G L	1100	s
	AAAATTA TCC AAAGGACCA CGGCGCTAG TGAGGAAGT ATCCAGCTA K L S K G T R A L S E E R I Q P I region B	1150	e
	TACTGCTTA TCTCTCTCC AAAACCTTA AGGACTAGG AGGCTCTT L A Y P H P K T L K Q L R G F L	1200	H
	GGCGATACAG GTTCTCTCCG AAAACCTTA CGGACTAGG CGCGCTACG G I T G F C R K Q I P R Y T P I A region A	1250	

53/69

FIG 47A



54/69

FIG 47B

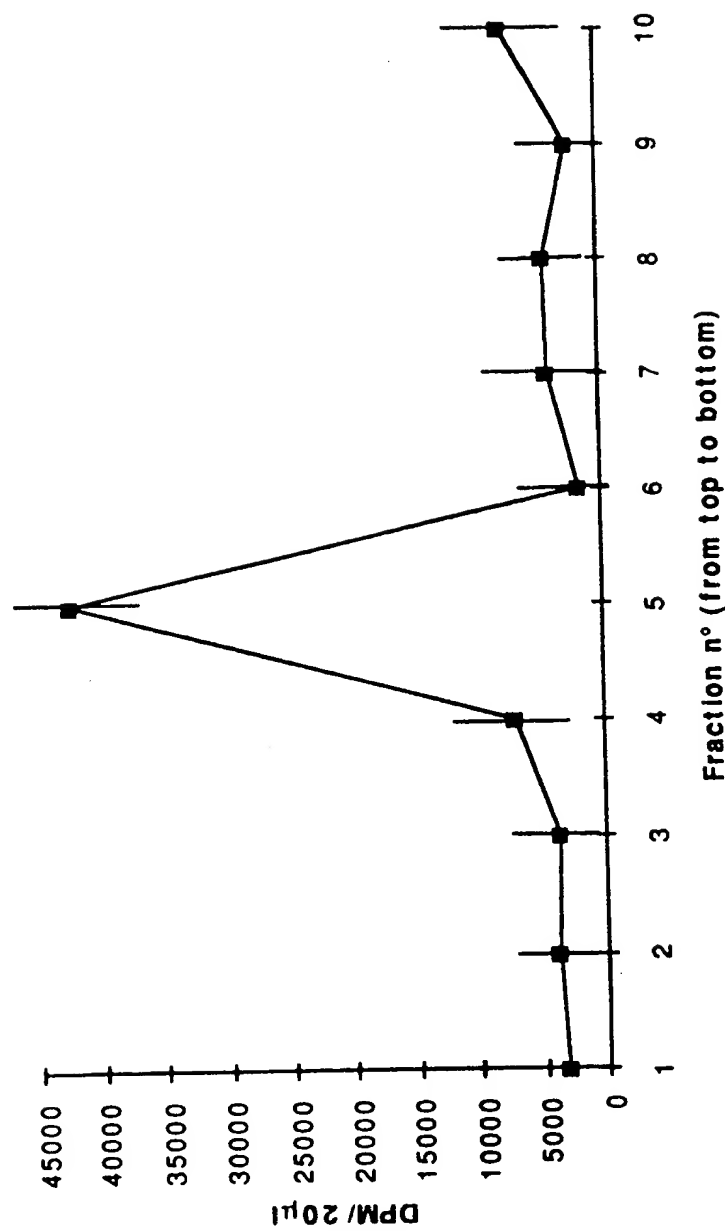


FIG 48A 55/69

	10	20	30	40	50	
<u>1234567890</u>	<u>1234567890</u>	<u>1234567890</u>	<u>1234567890</u>	<u>1234567890</u>	<u>1234567890</u>	
ATGATCCAGC	AGCAGGAACNG	AGGGTGCGCG	GGGCAAGCGC	CAGCCCCATGC		50
M I Q Q	Q D X	G C P	G Q A P	A H A		
CATCACCCCTC ACAGAGOCOC						100
I T L	T E P Q	V C L	T I E	G Q K G		
GTINACTGTCT CCTGGACACT GGCGGNGCCT TCTCAGTCIT ACTTTOCTGT						150
X C L	L D T	G G A F	S V L	L S C		
CCTGGACAAC TGTCCTCCAG ATCTGTCACT GTCCGAGGGG TCCTAGGACA						200
P G Q L	S S R	S V T	V R G V	L G Q		
GCCAGTCACT AGATACTTCT CCCAGCCACT AAGTTGTGAC TGGGGAACTT						250
P V T	R Y F S	Q P L	S C D	W G T L		
TACTCTTCCC ACACTGTTTT CTAATTATGC CTGAAAGCCC CACTCICITG						300
L F P	H A F	L I M P	E S P	T L L		
TTGGGGAGAG ACATTCTAGC AAAAGCAGGG GCCATTATAC ATGTGAATAT						350
L G R D	I L A	K A G	A I I H	V N I		
AGGAGAAGGA ACAACTGTIT GTTGCCCCCT GCTTGAGGAA GGAATTAATC						400
G E G	T T V C	C P L	L E E	G I N P		
CTGAAGTCGG GGCAACAGAA GGACAATATG GACAAGCAA GAATGCCCGT						450
E V R	A T E	G Q Y G	Q A K	N A R		
CCTGTTCAAG TTAAACTAAA GGATTCCACC TCCCTTCCCT ACCAAAGGCA						500
P V Q V	K L K	D S T	S F P Y	Q R Q		

## FIG 48B 56/69

10	20	30	40	50	
1234567890	1234567890	1234567890	1234567890	1234567890	550
GTACCCCCCTC	AGACCCGAGA	CCCAACAAGA	ACTCCAAAAG	ATTGTAAAGG	
Y P L	R P E T	Q Q E	L Q K	I V K D	
					600
ACCTAAAAGC	CCAAGGCCTA	GTAAAACCAA	GCAATAGCCC	TTGCAAGACT	
L K A	Q G L	V K P S	N S P	C K T	
					650
CCAATTTTAG	GAGTAAGGAA	ACCCAACGGA	CAGTGGAGGT	TAGTGCAAGA	
P I L G	V R K	P N G	Q W R L	V Q E	
					700
ACTCAGGATT	ATCAATGAGG	CTGTTGTCC	TCTATAACCA	GCTGTACCTA	
L R I	I N E A	V V P	L Y P	A V P N	
					750
ACCCTTATAC	AGTGCTTTC	CAAATACCG	AGGAAGCAGA	GTGGTTACA	
P Y T	V L S	Q I P E	E A E	W F T	
					800
GTCCTGGACC	TTAAGGATGC	CTTTTCTGC	ATCCTGTAC	GTCCTGACTC	
V L D L	K D A	F F C	I P V R	P D S	
					850
TCAATTCTTG	TTTGCCTTIG	AAGATCCTTT	GAACCCAACG	TCTCAACTCA	
Q F L	F A F E	D P L	N P T	S Q L T	
					900
CCTGGACTGT	TTTACCCCAA	GGGTCAGGG	ATAGCCCCCC	TCTATTGGC	
W T V	L P Q	G F R D	S P H	L F G	
					950
CAGGCATTAG	CCCAAGACIT	GAGTCATTIC	TCATAACCTGG	ACACTCTTGT	
Q A L A	Q D L	S Q F	S Y L D	T L V	
					1000
CCTTCAGTAC	ATGGATGATT	TACTTTAGT	CGCCCGTCA	GAAACCTTGT	
L Q Y	M D D L	L L V	A R S	E T L C	

## FIG 48C 57/69

10	20	30	40	50	
<u>1234567890</u>	<u>1234567890</u>	<u>1234567890</u>	<u>1234567890</u>	<u>1234567890</u>	
GCCATCAAGC	CACCCAAGAA	CTCTTAACIT	TCCTCACTAC	CTGTGGCTAC	1050
H Q A	T Q E	L L T F	L T T	C G Y	
AAGGTTTCCA AACCAAAGGC TOGGCTCTGC TCACAGGAGA TTAGATACTIN					1100
K V S K	P K A	R L C	S Q E I	R Y X	
AGGGCTAAAA TTATCCAAAG GCACCAGGGC OCTCAGTGAG GAACGTATCC					1150
G L K	L S K G	T R A	L S E	E R I Q	
AGCCTATACT GGCTTATCCT CATCCCCAAA CCTAAAGCA ACTAAGAGGG					1200
P I L	A Y P	H P K T	L K Q	L R G	
TTCCCTGGCA TAACAGGTTT CTGCCGAAAA CAGATTCCCA CGTACASCCC					1250
F L G I	T G F	C R K	Q I P R	Y X P	
AATAGCCAGA CCATTATATA CACTAATTAN GGAAACTCAG AAAGCCAATA					1300
I A R	P L Y T	L I X	E T Q	K A N T	
CCTATTTAGT AAGATGGACA CCTACAGAAG TGCTTTCCA GGCGCTAAAG					1350
Y L V	R W T	P T E V	A F Q	A L K	
AAGGCCCTAA CCCAAGCCCC AGTGTTCAGC TTGCCAACAG GGCAAGATTT					1400
K A L T	Q A P	V F S	L P T G	Q D F	
TTCTTTATAT GCCACAGAAA AAACAGGAAT AGCTCTAGGA GTCCCTACCG					1450
S L Y	A T E K	T G I	A L G	V L T Q	
AGGTCTCAGG GATGAGCTTG CAACCCGTGG TATACCTGAG TAAGGAAATT					1500
V S G	M S L	Q P V V	Y L S	K E I	

## FIG 48D

58/69

10	20	30	40	50	
<u>1234567890</u>	<u>1234567890</u>	<u>1234567890</u>	<u>1234567890</u>	<u>1234567890</u>	
GATGTAGTGG	CAAAGGGTG	GCCTCATNGT	TTATGGTAA	TGNGGCAGT	1550
D V V A	K G W	P H X	L W V M	X A V	
AGCAGTCINA	GTATCTGAAG	CAGTTAAAAT	AATACAGGGA	AGAGAICTIN	1600
A V X	V S E A	V K I	I Q G	R D L X	
CIGIGIGGAC	ATCTCATGAT	GTGAACGGCA	TACTSRCTGC	TAAAGGAGAC	1650
V W T	S H D	V N G I	L X A	K G D	
TTGIGGTTGT	CAGACAACCA	TTTACTTAAN	TAYCAGGCCYY	TATTACTTGA	1700
L W L S	D N H	L L X	Y Q A L	L L E	
AGAGCCAGTG	CTGNGACTGC	GCACTTGICC	AACCTCTAAA	CCCAAACCTTA	1750
E P V	L X L R	T C P	T L K	P K L M	
TGCTGCCAG	AAGGATCTTT	NTAGAGGTCC	CTTAGGCCAA	CCCTGACCIC	1800
L P R	R I F	X E V P	L A N	P D L	
AACTATATAT	ATACTGATGG	AAGTTGTTT	GTAGAAAAGG	GATTACAAAG	1850
N Y I Y	T D G	S S F	V E K G	L Q R	
CGNAGGATAT	NCCATAGGTG	TTAGTGATAA	ACCACTACTT	GAAAGTAAGC	1900
X G Y	X I G V	S D K	A V L	E S K P	
CTCTTCCCCC	CCAGGGACCA	GCGCCCCCGT	TAGCAGAACT	AGTGGCACTG	1950
L P P	Q G P	A P P L	A E L	V A L	
ACCCCCGGAG	CCTTAGAACT	TTGGAAAGGG	AGGAGGATAA	ATGTGTATAC	2000
T P R A	L E L	W K G	R R I N	V Y T	

59/69

## FIG 48E

10	20	30	40	50	
1234567890	1234567890	1234567890	1234567890	1234567890	
AGATAGCAAG	TATGCTTATC	TAATCCGAAA	TGCCCATGT	GCAATATGGA	2050
D S K	Y A Y L	I R N	A H V	A I W K	
<hr/>					
AAGAAAGGGA GTCCTAACCC TCTGGGGAA CCCCCATTAA ATACCACAAG					
E R E	F L T	S G G T	P I K	Y H K	2100
<hr/>					
TTAACATGG AGTTATTGCA CACAGTGCAA AAACCTCAAGG AGGTGGAAGT					
L I M E	L L H	T V Q	K L K E	V E V	2150
<hr/>					
CTTACACTGC CAAAGCCATC AGAAAAGGGA AAGACGGGAA GAGCAGCATA					
L H C	Q S H Q	K R E	R G E	E Q H K	2200
<hr/>					
AGTGGCTACA GAGGCAAGGA AAGACTAGCA GAAAGGAAAG AGAGAAAGAG					
W L Q	R Q G	K T S R	K E R	E K E	2250
<hr/>					
ACAGAAAGTC AGAGAGAGAG AGAGGAAGAG ACAGAGCACA AAGAGGGAGT					
T E S Q	R E R	E E E	T E H K	E G V	2300
<hr/>					
CAGAGAGAGA GAGAGACAGA GAGTCAGAGA GAAGGAAAGA GAGAGAGGAA					
R E R	E R Q R	V R E	K E R	E R G R	2350
<hr/>					
GAGACAAAGA ATGA					
D K E	.				2364

60/69  
FIG 49 A

Complement of 8/46-7 propre  
1 /46-7 propre  
Complement of c15 propre 46-7  
Consensus

Complement of 8/46-7 propre  
1 /46-7 propre  
Complement of c15 propre 46-7  
Consensus

Complement of 8/46-7 propre  
1 /46-7 propre  
Complement of c15 propre 46-7  
Consensus

Complement of 8/46-7 propre  
1 /46-7 propre  
Complement of c15 propre 46-7  
Consensus

Complement of 8/46-7 propre  
1 /46-7 propre  
Complement of c15 propre 46-7  
Consensus

Complement of 8/46-7 propre  
1 /46-7 propre  
Complement of c15 propre 46-7  
Consensus

Complement of 8/46-7 propre  
1 /46-7 propre  
Complement of c15 propre 46-7  
Consensus

Complement of 8/46-7 propre  
1 /46-7 propre  
Complement of c15 propre 46-7  
Consensus

Complement of 8/46-7 propre  
1 /46-7 propre  
Complement of c15 propre 46-7  
Consensus

GACTTGAGCC AGTCCTCATA CCTGGACATT CTTGTTCTTC AGTATGGGA	50
GACTTGAGCC AGTCCTCATA CCTGGACATT CTTGTTCTTC AGTATGGGA	50
GACTTGAGCC AGTCCTCATA CCTGGACATT CTTGTTCTTC AGTATGGGA	50
GACTTGAGCC AGTCCTCATA CCTGGACATT CTTGTTCTTC AGTATGGGA	50

TGA[TTAATT ATAGCCACCC ATTCAAGAAC CTTGTGGCAT CAAGCCACCC	100
TGA[TTAATT ATAGCCACCC ATTCAAGAAC CTTGTGGCAT CAAGCCACCC	100
TGA[TTAATT ATAGCCACCC ATTCAAGAAC CTTGTGGCAT CAAGCCACCC	100
TGA[TTAATT ATAGCCACCC ATTCAAGAAC CTTGTGGCAT CAAGCCACCC	100

AAG[GCTCTT AAATTCCTC GCTACCTGTG GCTCCAAACA AAGGGCTCAG	150
AAG[GCTCTT AAATTCCTC GCTACCTGTG GCTCCAAACA AAGGGCTCAG	150
AAG[GCTCTT AAATTCCTC GCTACCTGTG GCTCCAAACA AAGGGCTCAG	150
AAG[GCTCTT AAATTCCTC GCTACCTGTG GCTCCAAACA AAGGGCTCAG	150

CTCTGCTCAC ACCAGGTTAA ATACTTAGGG CTAAAATTAT CCAAAGTCAG	200
CTCTGCTCAC ACCAGGTTAA ATACTTAGGG CTAAAATTAT CCAAAGTCAG	200
CTCTGCTCAC ACCAGGTTAA ATACTTAGGG CTAAAATTAT CCAAAGTCAG	200
CTCTGCTCAC ACCAGGTTAA ATACTTAGGG CTAAAATTAT CCAAAGTCAG	200

CAGGGCCCTC AGAGAGGAAC GTATCCAGCG TATACTGGAT TATCCCATC	250
CAGGGCCCTC AGAGAGGAAC GTATCCAGCG TATACTGGAT TATCCCATC	250
CAGGGCCCTC AGAGAGGAAC GTATCCAGCG TATACTGGAT TATCCCATC	250
CAGGGCCCTC AGAGAGGAAC GTATCCAGCG TATACTGGAT TATCCCATC	250

CCAAACCTT AAAGCAACTA AGA[GGTTCC TTGGCATATC AGCCTTCCTGC	300
CCAAACCTT AAAGCAACTA AGA[GGTTCC TTGGCATATC AGCCTTCCTGC	300
CCAAACCTT AAAGCAACTA AGA[GGTTCC TTGGCATATC AGCCTTCCTGC	300
CCAAACCTT AAAGCAACTA AGA[GGTTCC TTGGCATATC AGCCTTCCTGC	300

CGAATATGGA TTCCC[GATA CAG[GAAATA GCCAGGCCAT TATGTACATT	350
CGAATATGGA TTCCC[GATA CAG[GAAATA GCCAGGCCAT TATGTACATT	350
CGAATATGGA TTCCC[GATA CAG[GAAATA GCCAGGCCAT TATGTACATT	350
CGAATATGGA TTCCC[GATA CAG[GAAATA GCCAGGCCAT TATGTACATT	350

AG[TAAGGAA ACTCAGAAAG CCAATACCCA TATAGTAAGA TGGACACCTG	400
AG[TAAGGAA ACTCAGAAAG CCAATACCCA TATAGTAAGA TGGACACCTG	400
AG[TAAGGAA ACTCAGAAAG CCAATACCCA TATAGTAAGA TGGACACCTG	400
AG[TAAGGAA ACTCAGAAAG CCAATACCCA TATAGTAAGA TGGACACCTG	400

ATACAGAAAGT GGCTTTCCAG GCCCTAAAG	429

## FIG 49B

Trans of 1 /46-7 pr	DLSQSSYLDI	LVIQYEDDLI	IATHSETLWH	QATO	LLNFL	ATCGSKORAQ	50
Trans of Complement-2(8)	DLSQSSYLDI	LVIQYFDDLI	IATHSETLWH	QATO	LLNFL	ATCGSKORAQ	50
Trans of Complement(5)	DLSQSSYLDI	LVIQYEDDLI	IATHSETLWH	QATO	LLNFL	ATCGSKORAQ	50
Consensus	DLSQSSYLDI	LVIQYEDDLI	IATHSETLWH	QATO	LLNFL	ATCGSKORAQ	50
Trans of 1 /46-7 pr	LCSQVKYLG	LKLSKVIRAL	REERIQRILIA	YPHE	IIKQL	REFLGIISAFQ	100
Trans of Complement-2	LCSQVKYLG	LKLSKVIRAL	REERIQRILD	YPHE	IIKQL	REFLGIISAFQ	100
Trans of Complement	LCSQVKYLG	LKLSKVIRAL	REERIQRILIA	YPHE	IIKQL	REFLGIISAFQ	100
Consensus	LCSQVKYLG	LKLSKVIRAL	REERIQRILIA	YPHE	IIKQL	REFLGIISAFQ	100
Trans of 1 /46-7 pr	RIWIPSYSEI	ARPLCTLKE	TOKANTHIVR	WTPETEVAFQ	ALK		143
Trans of Complement-2	RIWIPSYSEI	ARPLCTLKE	TOKANTHIVR	WTPETEVAFQ	ALK		143
Trans of Complement	RIWIPSYSEI	ARPLCTLKE	TOKANTHIVR	WTPETEVAFQ	ALK		143
Consensus	RIWIPSYSEI	ARPLCTLKE	TOKANTHIVR	WTPETEVAFQ	ALK		143

## FIG 50B

Trans of c143 propr	DLSQSSYLDX	LVLRYMDDL	LATHSETLCH	QATO	LLNFL	ATCGYKVSKP	50
Trans of 42/68-1 pr	DLSQSSYLDT	LVLRYMDDL	LATHSETLCH	QATO	LLNFL	ATCGYKVSKP	50
Trans of 41/68-1 pr	DLSQSSYLDT	LVLRYMDDL	LATHSETLCH	QATO	LLNFL	ATCGYKVSKP	50
Consensus	DLSQSSYLDT	LVLRYMDDL	LATHSETLCH	QATO	LLNFL	ATCGYKVSKP	50
Trans of c143 propr	KAQLCSQQVK	YLGLKLSKGT	RTLSEERIQP	ILGYPHPKTL	KQLTAFLGIT		100
Trans of 42/68-1 pr	KAQLCSQQVK	YLGLKLSKGT	RTLSEERIQP	ILGYPHPKTL	KQLTAFLGIT		100
Trans of 41/68-1 pr	KAQLCSQQVK	YLGLKLSKGT	RTLSEERIQP	ILGYPHPKTL	KQLTAFLGIT		100
Consensus	KAQLCSQQVK	YLGLKLSKGT	RTLSEERIQP	ILGYPHPKTL	KQLTAFLGIT		100
Trans of c143 propr	GFCQIWIPRY	SKMARPLNTR	IKETOKANITH	LVRWISAEAV	AFQALK		146
Trans of 42/68-1 pr	GFCQIWIPRY	SKMARPLNTR	IKETOKAETH	LVRWISAEAV	AFQALK		146
Trans of 41/68-1 pr	GFCQIWIPRY	SKMARPLNTR	IKETOKANITH	LVRWISAEAV	AFQALK		146
Consensus	GFCQIWIPRY	SKMARPLNTR	IKETOKANITH	LVRWISAEAV	AFQALK		146

62/69

## FIG 50 A

41/68-1 propre	GACTTGAGCC AGTGTTCATA CCTGGACACT CTTGTCCTTC GGTACATGGG	50
c143 propre 68-1	GACTTGAGCC AGTGTTCATA CCTGGACACT CTTGTCCTTC GGTACATGGG	50
42/68-1 propre	GACTTGAGCC AGTGTTCATA CCTGGACACT CTTGTCCTTC GGTACATGGG	50
Consensus	GACTTGAGCC AGTGTTCATA CCTGGACACT CTTGTCCTTC GGTACATGGG	50
41/68-1 propre	TGATTTACTT TTAGGCCACCC ATTCAAGAAAC CTTGTGCCAT CAAGCCACCC	100
c143 propre 68-1	TGATTTACTT TTAGGCCACCC ATTCAAGAAAC CTTGTGCCAT CAAGCCACCC	100
42/68-1 propre	TGATTTACTT TTAGGCCACCC ATTCAAGAAAC CTTGTGCCAT CAAGCCACCC	100
Consensus	TGATTTACTT TTAGGCCACCC ATTCAAGAAAC CTTGTGCCAT CAAGCCACCC	100
41/68-1 propre	AAGCACTCTT AAATTTCCCT GCTACCTGTG GCTACAAGGT TTCCAAACCA	150
c143 propre 68-1	AAGCACTCTT AAATTTCCCT GCTACCTGTG GCTACAAGGT TTCCAAACCA	150
42/68-1 propre	AAGCACTCTT AAATTTCCCT GCTACCTGTG GCTACAAGGT TTCCAAACCA	150
Consensus	AAGCACTCTT AAATTTCCCT GCTACCTGTG GCTACAAGGT TTCCAAACCA	150
41/68-1 propre	AAGGCTCAGC TCTGCTCACCA GCAGGTTAAA TACTTGGGC TAAAATTATC	200
c143 propre 68-1	AAGGCTCAGC TCTGCTCACCA GCAGGTTAAA TACTTGGGC TAAAATTATC	200
42/68-1 propre	AAGGCTCAGC TCTGCTCACCA GCAGGTTAAA TACTTGGGC TAAAATTATC	200
Consensus	AAGGCTCAGC TCTGCTCACCA GCAGGTTAAA TACTTGGGC TAAAATTATC	200
41/68-1 propre	CAAAGGCACC AGAACCCCTCA GTGAGGAACG TATCCAGCCT ATACTGGGT	250
c143 propre 68-1	CAAAGGCACC AGAACCCCTCA GTGAGGAACG TATCCAGCCT ATACTGGGT	250
42/68-1 propre	CAAAGGCACC AGAACCCCTCA GTGAGGAACG TATCCAGCCT ATACTGGGT	250
Consensus	CAAAGGCACC AGAACCCCTCA GTGAGGAACG TATCCAGCCT ATACTGGGT	250
41/68-1 propre	ATCCTCATCC CAAAACCTTA AAGCAACTAA CAGCGTTCCCT TGGCATAACA	300
c143 propre 68-1	ATCCTCATCC CAAAACCTTA AAGCAACTAA CAGCGTTCCCT TGGCATAACA	300
42/68-1 propre	ATCCTCATCC CAAAACCTTA AAGCAACTAA CAGCGTTCCCT TGGCATAACA	300
Consensus	ATCCTCATCC CAAAACCTTA AAGCAACTAA CAGCGTTCCCT TGGCATAACA	300
41/68-1 propre	GGTTTCTGCC AAATATGGAT TCCCAGGTAC AGCAAGTAG CCAGACCATT	350
c143 propre 68-1	GGTTTCTGCC AAATATGGAT TCCCAGGTAC AGCAAGTAG CCAGACCATT	350
42/68-1 propre	GGTTTCTGCC AAATATGGAT TCCCAGGTAC AGCAAGTAG CCAGACCATT	350
Consensus	GGTTTCTGCC AAATATGGAT TCCCAGGTAC AGCAAGTAG CCAGACCATT	350
41/68-1 propre	AAATACACGA ATTAAGGAAA CTCAAAAAGC CAGTACCCAT TTAGTAAGAT	400
c143 propre 68-1	AAATACACGA ATTAAGGAAA CTCAAAAAGC CAGTACCCAT TTAGTAAGAT	400
42/68-1 propre	AAATACACGA ATTAAGGAAA CTCAAAAAGC CAGTACCCAT TTAGTAAGAT	400
Consensus	AAATACACGA ATTAAGGAAA CTCAAAAAGC CAGTACCCAT TTAGTAAGAT	400
41/68-1 propre	GGACACCTGA AGCAGAAGTG GCTTTCCAGG CCCTAAAG	438
c143 propre 68-1	GGACACCTGA AGCAGAAGTG GCTTTCCAGG CCCTAAAG	438
42/68-1 propre	GGACACCTGA AGCAGAAGTG GCTTTCCAGG CCCTAAAG	438
Consensus	GGACACCTGA AGCAGAAGTG GCTTTCCAGG CCCTAAAG	438

## FIG 51A

MSRV pol cons ADN 1,5,8	ATTATGCCCTG AAAGCCCCAC TCCCTTGTAA GGGAGAGACA TTTTAGCAAA	50
Consensus	ATTATGCCCTG AAAGCCCCAC TCCCTTGTAA GGGAGAGACA TTTTAGCAAA	50
MSRV pol cons ADN 1,5,8	AGCAGGGGCC ATTATACACC TGAACATAGG AAAAGGAATA CCCATTTGCT	100
Consensus	AGCAGGGGCC ATTATACACC TGAACATAGG AAAAGGAATA CCCATTTGCT	100
MSRV pol cons ADN 1,5,8	GTCCTCTGCT TGAGGAAGGA ATTAATCCTG AAGTCTGGC AATAGAAGGA	150
Consensus	GTCCTCTGCT TGAGGAAGGA ATTAATCCTG AAGTCTGGC AATAGAAGGA	150
MSRV pol cons ADN 1,5,8	CAATATGGAC AAGCAAAGAA TGCCCGTCCT GTTCAAGTTA AACTAAAGGA	200
Consensus	CAATATGGAC AAGCAAAGAA TGCCCGTCCT GTTCAAGTTA AACTAAAGGA	200
MSRV pol cons ADN 1,5,8	TTCCTGCCTCC TTTCCTTACCC AAAGGAAGTA CCCTCTTAGA CCCCAGGGCCC	250
Consensus	TTCCTGCCTCC TTTCCTTACCC AAAGGAAGTA CCCTCTTAGA CCCCAGGGCCC	250
MSRV pol cons ADN 1,5,8	TACAAGGANC TCAAAAGATT GTTAAGGACC TAAAAGCCC AGGCCTAGTA	300
Consensus	TACAAGGANC TCAAAAGATT GTTAAGGACC TAAAAGCCC AGGCCTAGTA	300
MSRV pol cons ADN 1,5,8	AAACCATGCA GTAGCCCCTG CAATACTCCA ATTTTAGGAG TAAGGAAACC	350
Consensus	AAACCATGCA GTAGCCCCTG CAATACTCCA ATTTTAGGAG TAAGGAAACC	350
MSRV pol cons ADN 1,5,8	CAACGGACAG TGGAGGTAG TGCAAGATCT CAGGATTATT AATGAGGCTG	400
Consensus	CAACGGACAG TGGAGGTAG TGCAAGATCT CAGGATTATT AATGAGGCTG	400
MSRV pol cons ADN 1,5,8	TTTTTCCCTCT ATACCCAGCT GTATCTAGCC CTTATACTCT GCTTTCCCTA	450
Consensus	TTTTTCCCTCT ATACCCAGCT GTATCTAGCC CTTATACTCT GCTTTCCCTA	450
MSRV pol cons ADN 1,5,8	ATACCAAGAGG AAGCAGAGTG GTTTACAGTC CTGGACCTTA AGGATGCCTT	500
Consensus	ATACCAAGAGG AAGCAGAGTG GTTTACAGTC CTGGACCTTA AGGATGCCTT	500
MSRV pol cons ADN 1,5,8	TTTCTGCATC CCTGTACGTC CTGACTCTCA ATTCTGTGTT GCCTTGTAAAG	550
Consensus	TTTCTGCATC CCTGTACGTC CTGACTCTCA ATTCTGTGTT GCCTTGTAAAG	550
MSRV pol cons ADN 1,5,8	ATCCTTTGAA CCCAACGTCT CAACTCACCT GGACTGTGTT ACCCCAAGGG	600
Consensus	ATCCTTTGAA CCCAACGTCT CAACTCACCT GGACTGTGTT ACCCCAAGGG	600
MSRV pol cons ADN 1,5,8	TTCAGGGATA GCCCCCATCT ATTGGCCAG GCATTAGCCC A[REDACTED]GAG [REDACTED]GAG	650
Consensus	TTCAGGGATA GCCCCCATCT ATTGGCCAG GCATTAGCCC A[REDACTED]GAG	650
MSRV pol cons ADN 1,5,8	[REDACTED]TCA TACCTGGACCA [REDACTED]CTCTGCTC [REDACTED]CAGTACGIG GATGAA[TTC] [REDACTED]CTCA TACCTGGACCA [REDACTED]CTCTGCTC [REDACTED]CAGTACGIG GATGAA[TTC] [REDACTED]CTCA TACCTGGACCA [REDACTED]CTCTGCTC [REDACTED]CAGTACGIG GATGAA[TTC]	700
Consensus	[REDACTED]TCA TACCTGGACCA [REDACTED]CTCTGCTC [REDACTED]CAGTACGIG GATGAA[TTC] [REDACTED]CTCA TACCTGGACCA [REDACTED]CTCTGCTC [REDACTED]CAGTACGIG GATGAA[TTC] [REDACTED]CTCA TACCTGGACCA [REDACTED]CTCTGCTC [REDACTED]CAGTACGIG GATGAA[TTC]	700
MSRV pol cons ADN 1,5,8	[REDACTED]TACCTGGACCA [REDACTED]CTCTGCTC [REDACTED]CAGTACGIG GATGAA[TTC] [REDACTED]TACCTGGACCA [REDACTED]CTCTGCTC [REDACTED]CAGTACGIG GATGAA[TTC] [REDACTED]TACCTGGACCA [REDACTED]CTCTGCTC [REDACTED]CAGTACGIG GATGAA[TTC]	750
Consensus	[REDACTED]TACCTGGACCA [REDACTED]CTCTGCTC [REDACTED]CAGTACGIG GATGAA[TTC] [REDACTED]TACCTGGACCA [REDACTED]CTCTGCTC [REDACTED]CAGTACGIG GATGAA[TTC] [REDACTED]TACCTGGACCA [REDACTED]CTCTGCTC [REDACTED]CAGTACGIG GATGAA[TTC]	750
MSRV pol cons ADN 1,5,8	[REDACTED]TACCTGGACCA [REDACTED]CTCTGCTC [REDACTED]CAGTACGIG GATGAA[TTC] [REDACTED]TACCTGGACCA [REDACTED]CTCTGCTC [REDACTED]CAGTACGIG GATGAA[TTC] [REDACTED]TACCTGGACCA [REDACTED]CTCTGCTC [REDACTED]CAGTACGIG GATGAA[TTC]	800
Consensus	[REDACTED]TACCTGGACCA [REDACTED]CTCTGCTC [REDACTED]CAGTACGIG GATGAA[TTC] [REDACTED]TACCTGGACCA [REDACTED]CTCTGCTC [REDACTED]CAGTACGIG GATGAA[TTC] [REDACTED]TACCTGGACCA [REDACTED]CTCTGCTC [REDACTED]CAGTACGIG GATGAA[TTC]	800

## FIG 51A (cont.)

MSRV pol cons ADN 1,5,8 Consensus	GCTCTGCTCA CASSAGCTTA TATACTTAGG GCTAAAATTA TCCAAAGCCT SCTCTGCTCA CASSAGCTTA TATACTTAGG GCTAAAATTA TCCAAAGCCT SCTCTGCTCA CASSAGCTTA TATACTTAGG GCTAAAATTA TCCAAAGCCT	850 199 850
MSRV pol cons ADN 1,5,8 Consensus	CCAGGGCCCT CAGGAGGA CGTATCCAGC TATACTGGC TTATCCCAT CCAGGGCCCT CAGGAGGA CGTATCCAGC TATACTGGM TTATCCCAT CCAGGGCCCT CAGGAGGA CGTATCCAGC TATACTGGM TTATCCCAT	900 249 900
MSRV pol cons ADN 1,5,8 Consensus	CCCANAACCC TAAAGCAACT AAGAGGGTC CTTGCCATAA CAGGTTCTGC CCCANAACCC TAAAGCAACT AAGAGGGTC CTTGCCATAW CAGGTTCTGC CCCANAACCC TAAAGCAACT AAGAGGGTC CTTGCCATAW CAGGTTCTGC	950 299 950
MSRV pol cons ADN 1,5,8 Consensus	CGGAATACCG ATTCCCGT ACACCCGAT AGCCAGCCA TTATCTACAT CGGAATACCG ATTCCCGT ACAGYGAAT AGCCAGCCA TTATCTACAT CGGAATACCG ATTCCCGT ACASYSMAAT AGCCAGCCA TTATCTACAT	1000 349 1000
MSRV pol cons ADN 1,5,8 Consensus	TAAITTAAGGA AACTCAGAAA GCCAATACCC ATTTAGTAAG ATGGACACCT TADYTARGGA AACTCAGAAA GCCAATACCC ATTTAGTAAG ATGGACACCT TADYTARGGA AACTCAGAAA GCCAATACCC ATTTAGTAAG ATGGACACCT	1050 399 1050
MSRV pol cons ADN 1,5,8 Consensus	-- ACAGAAG TGGCTTCGA GGCCCTAAAG AAGGCCCTAA CCCAAGCCCC GARACAGAAG TGGCTTCGA GGCCCTAAAG ----- GARACAGAAG TGGCTTCGA GGCCCTAAAG AAGGCCCTAA CCCAAGCCCC	1097 429 1100
MSRV pol cons ADN 1,5,8 Consensus	AGTGTTCAAGC TTGCCAACAG GGCAAGATT TTCTTTATAT GCCACAGAAA ----- AGTGTTCAAGC TTGCCAACAG GGCAAGATT TTCTTTATAT GCCACAGAAA	1147 429 1150
MSRV pol cons ADN 1,5,8 Consensus	AAACAGGAAT AGCTCTAGGA GTCTTACGC AGGTCTCAGG GATGAGCTTG ----- AAACAGGAAT AGCTCTAGGA GTCTTACGC AGGTCTCAGG GATGAGCTTG	1197 429 1200
MSRV pol cons ADN 1,5,8 Consensus	CAACCCGTGG TATACCTGAG TAAGGAAATT GATGTAGTGG CAAAGGGTTG ----- CAACCCGTGG TATACCTGAG TAAGGAAATT GATGTAGTGG CAAAGGGTTG	1247 429 1250
MSRV pol cons ADN 1,5,8 Consensus	GCCTCATTGT TTATGGTAA TGGCGGCAGT AGCAGTCTTA GTATCTGAAG ----- GCCTCATTGT TTATGGTAA TGGCGGCAGT AGCAGTCTTA GTATCTGAAG	1297 429 1300
MSRV pol cons ADN 1,5,8 Consensus	CAGTTAAAAT AATACAGGGA AGAGATCTTA CTGTGTGGAC ATCTCATGAT ----- CAGTTAAAAT AATACAGGGA AGAGATCTTA CTGTGTGGAC ATCTCATGAT	1347 429 1350
MSRV pol cons ADN 1,5,8 Consensus	GTGAACGGCA TACTCACTGC TAAAGGAGAC TTGTGGTTGT CAGACAACCA ----- GTGAACGGCA TACTCACTGC TAAAGGAGAC TTGTGGTTGT CAGACAACCA	1397 429 1400
MSRV pol cons ADN 1,5,8 Consensus	TTTACTTAAT TATCAGGCTC TATTAATTGA AGAGCCAGTG CTGAGACTGC ----- TTTACTTAAT TATCAGGCTC TATTAATTGA AGAGCCAGTG CTGAGACTGC	1447 429 1450
MSRV pol cons ADN 1,5,8 Consensus	GCACTTGTGC AACTCTAAA CCCGCCACAT TTCTTCCAGA CAATGAAGAA ----- GCACTTGTGC AACTCTAAA CCCGCCACAT TTCTTCCAGA CAATGAAGAA	1497 429 1500
MSRV pol cons ADN 1,5,8 Consensus	AAGATAGAAC ATAACGTCA ACAAGTAATT GCTCAAACCT ATGCTGCTCG ----- AAGATAGAAC ATAACGTCA ACAAGTAATT GCTCAAACCT ATGCTGCTCG	1547 429 1550
MSRV pol cons ADN 1,5,8 Consensus	AGGGGACCTT CTAGAGGTTT CCTTGACTGA TCCCGACCTC AACTTGTATA ----- AGGGGACCTT CTAGAGGTTT CCTTGACTGA TCCCGACCTC AACTTGTATA	1597 429 1600

65/69

## FIG 51B

Trans of MSRV pol cons prot 1,5,8	IMPESPTPLL GRDILAKAGA IIHLNIGKGI PICCPLLEEG INPEVWAIEG	50
Consensus	.....	50
Trans of MSRV pol cons prot 1,5,8	QYGQAKNARP VQVKLKDAS FPYQRKYPLR PEALQGXQKI VKDLKAQGLV	100
Consensus	.....	100
Trans of MSRV pol cons prot 1,5,8	KPCSSPCNTP ILGVRKPNGQ WRLVQDLRII NEAVFPLYPA VSSPYTLLSL	150
Consensus	.....	150
Trans of MSRV pol cons prot 1,5,8	IPEEEAEWFTV LDLKDAFFCI PVRPDQSFLF AFEDPLNPTS QLTWIVLPQG	200
Consensus	.....	200
Trans of MSRV pol cons prot 1,5,8	FRDSPHLFGQ ALAQ[DLSQES YLDILVLQYV [DDLI[LVARSE T[HQATOEL	250
Consensus	[DLSQES YLDILVLQYV [DDLI[LATHSE T[HQATOEL [DLSQES YLDILVLQYV [DDLI[LVARSE T[HQATOEL	36
Trans of MSRV pol cons prot 1,5,8	.....	250
Consensus	.....	
Trans of MSRV pol cons prot 1,5,8	[INFLATCGSK VSKE[KANLCS GEIRYLGLKL SK[TRALSEE RIC[TLAYPH [INFLATCGSK --KA[KANLCS GQVYLGLKL SK[TRALSEE RIC[TLAYPH [INFLATCGSK ...KA[KANLCS G...YLGLKL SK[TRALSEE RIC[TLAYPH	300
Consensus	.....	83
Trans of MSRV pol cons prot 1,5,8	PKTLKQLRGF LGIT[PCRKO IPRYTHIARP LMTLIRETQK ANIYLVRWTP PKTLKQLRGF LGIT[PCRKW IPRYSHIARP LMTLKKETQK ANIYLVRWTP PKTLKQLRGF LGIT[PCR... IPRY... IARP L...L... ETQK ANI... VRWTP	350
Consensus	.....	133
Trans of MSRV pol cons prot 1,5,8	TEVAEQALK KALTQAPVFS LPTGQDFSLY ATEKTGIALG VLTQVSGMSL	399
Consensus	TEVAEQALK	143
Trans of MSRV pol cons prot 1,5,8	.....	400
Trans of MSRV pol cons prot 1,5,8	QPVVYLSKEI DVVAKGPHC LWVMAAVAVL VSEAVKIIQG RDLTWWTSHD	449
Consensus	.....	143
Trans of MSRV pol cons prot 1,5,8	.....	450
Trans of MSRV pol cons prot 1,5,8	VNGILTAKGD LWLSDNHILN YQALLLEEPV LRLRTCATLK PATFLPDNEE	499
Consensus	.....	143
Trans of MSRV pol cons prot 1,5,8	.....	500
Trans of MSRV pol cons prot 1,5,8	KIEHNCQQVI AQTYAARGDL LEVPLTDPL NLYTDGSSLA EKGLRKAGYA	549
Consensus	.....	143
Trans of MSRV pol cons prot 1,5,8	.....	550
Trans of MSRV pol cons prot 1,5,8	VISDNGILES NRLTPGTSAH LAELIAITWA LELGEKGKVN IYSDSKYAYL	599
Consensus	.....	143
Trans of MSRV pol cons prot 1,5,8	.....	600
Trans of MSRV pol cons prot 1,5,8	VLHAAAIWR EREFLTSEG T PINHQEAIRR LLLAVQKPKE VAVLHCQGHQ	649
Consensus	.....	143
Trans of MSRV pol cons prot 1,5,8	.....	650
Trans of MSRV pol cons prot 1,5,8	EEEEEREIEGN RQADIEAKKA ARQDSPLEML IEGP	683
Consensus	.....	143
Trans of MSRV pol cons prot 1,5,8	.....	684
Consensus	.....	

66/69

## FIG 52 A

MSRV pol cons ADN 41,42,43	ATATATGCCTG AAAGCCCCAC TCCCTTGTAA GGGAGAGACA TTTTAGCAAA	50
Consensus	-----	
MSRV pol cons ADN 41,42,43	ATTATGCCTG AAAGCCCCAC TCCCTTGTAA GGGAGAGACA TTTTAGCAAA	50
Consensus	-----	
MSRV pol cons ADN 41,42,43	AGCAGGGGCC ATTATACACC TGAACATAGG AAAAGGAATA CCCATTTGCT	100
Consensus	-----	
MSRV pol cons ADN 41,42,43	AGCAGGGGCC ATTATACACC TGAACATAGG AAAAGGAATA CCCATTTGCT	100
Consensus	-----	
MSRV pol cons ADN 41,42,43	GTCCCCCTGCT TGAGGAAGGA ATTAATCCTG AAGTCTGGC AATAGAAGGA	150
Consensus	-----	
MSRV pol cons ADN 41,42,43	GTCCCCCTGCT TGAGGAAGGA ATTAATCCTG AAGTCTGGC AATAGAAGGA	150
Consensus	-----	
MSRV pol cons ADN 41,42,43	CAATATGGAC AAGCAAAGAA TGCCCGTCCT GTTCAAGTTA AACTAAAGGA	200
Consensus	-----	
MSRV pol cons ADN 41,42,43	CAATATGGAC AAGCAAAGAA TGCCCGTCCT GTTCAAGTTA AACTAAAGGA	200
Consensus	-----	
MSRV pol cons ADN 41,42,43	TTCTGCCTCC TTTCCTTACCC AAAGGAAGTA CCCTCTTAGA CCCGAGGCC	250
Consensus	-----	
MSRV pol cons ADN 41,42,43	TTCTGCCTCC TTTCCTTACCC AAAGGAAGTA CCCTCTTAGA CCCGAGGCC	250
Consensus	-----	
MSRV pol cons ADN 41,42,43	TACAAGGANC TCAAAAGATT GTTAAGGACC TAAAAGCCCA AGGCCTAGTA	300
Consensus	-----	
MSRV pol cons ADN 41,42,43	TACAAGGANC TCAAAAGATT GTTAAGGACC TAAAAGCCCA AGGCCTAGTA	300
Consensus	-----	
MSRV pol cons ADN 41,42,43	AAACCATGCA GTAGCCCTTG CAATACTCCA ATTTTGTAGG TAAGGAAACC	350
Consensus	-----	
MSRV pol cons ADN 41,42,43	AAACCATGCA GTAGCCCTTG CAATACTCCA ATTTTGTAGG TAAGGAAACC	350
Consensus	-----	
MSRV pol cons ADN 41,42,43	CAACGGACAG TGGAGGTAG TGCAAGATCT CAGGATTATT AATGAGGCTG	400
Consensus	-----	
MSRV pol cons ADN 41,42,43	CAACGGACAG TGGAGGTAG TGCAAGATCT CAGGATTATT AATGAGGCTG	400
Consensus	-----	
MSRV pol cons ADN 41,42,43	TTTTTCTCT ATACCCAGCT GTATCTAGCC CTTATACTCT GCTTTCCCTA	450
Consensus	-----	
MSRV pol cons ADN 41,42,43	TTTTTCTCT ATACCCAGCT GTATCTAGCC CTTATACTCT GCTTTCCCTA	450
Consensus	-----	
MSRV pol cons ADN 41,42,43	ATACCAGAGG AAGCAGAGTG GTTACAGTC CTGGACCTTA AGGATGCCTT	500
Consensus	-----	
MSRV pol cons ADN 41,42,43	ATACCAGAGG AAGCAGAGTG GTTACAGTC CTGGACCTTA AGGATGCCTT	500
Consensus	-----	
MSRV pol cons ADN 41,42,43	TTTCTGCATC CCTGTACGTC CTGACTCTCA ATTCTTGTIT GCCTTTGAAG	550
Consensus	-----	
MSRV pol cons ADN 41,42,43	TTTCTGCATC CCTGTACGTC CTGACTCTCA ATTCTTGTIT GCCTTTGAAG	550
Consensus	-----	
MSRV pol cons ADN 41,42,43	ATCCTTTGAA CCCAACGTCT CAACTCACCT GGACTGTGTTT ACCCCAAGGG	600
Consensus	-----	
MSRV pol cons ADN 41,42,43	ATCCTTTGAA CCCAACGTCT CAACTCACCT GGACTGTGTTT ACCCCAAGGG	600
Consensus	-----	
MSRV pol cons ADN 41,42,43	TTCAGGGATA GCCCCCACATCT ATTGGCCAG GCATTAGCCC AAGACTTGTAG	650
Consensus	-----	
MSRV pol cons ADN 41,42,43	TTCAGGGATA GCCCCCACATCT ATTGGCCAG GCATTAGCCC AAGACTTGTAG	650
Consensus	-----	
MSRV pol cons ADN 41,42,43	TCACTTGTCA TACCTGGACCA TCTTGTCTCT TCTGTACTTG GATGATTTCAC	700
Consensus	-----	
MSRV pol cons ADN 41,42,43	CCAGCTCTCA TACCTGGACCA TCTTGTCTCT TCTGTACTTG GATGATTTCAC	58
Consensus	-----	
MSRV pol cons ADN 41,42,43	TCAGTGTCA TACCTGGACCA TCTTGTCTCT TCTGTACTTG GATGATTTCAC	700
Consensus	-----	
MSRV pol cons ADN 41,42,43	TTTTAGCTTC CCGTTTCAGAA ACCTTGTGCC ATCAAGCCAC CCAAGGTACTC	750
Consensus	-----	
MSRV pol cons ADN 41,42,43	TTTTAGCTTC CCGTTTCAGAA ACCTTGTGCC ATCAAGCCAC CCAAGGTACTC	108
Consensus	-----	
MSRV pol cons ADN 41,42,43	TTTTAGCTTC CCGTTTCAGAA ACCTTGTGCC ATCAAGCCAC CCAAGGTACTC	750
Consensus	-----	
MSRV pol cons ADN 41,42,43	TTAACTTTCC TCACCTTG TGCTACAAG GTTTCAAAC CAAAGGTCTG	800
Consensus	-----	
MSRV pol cons ADN 41,42,43	TTAACTTTCC TTGCTACCTG TGCTACAAG GTTTCAAAC CAAAGGTCTG	158
Consensus	-----	
MSRV pol cons ADN 41,42,43	TTAACTTTCC TTGCTACCTG TGCTACAAG GTTTCAAAC CAAAGGTCTG	800
Consensus	-----	

67165

FIG 52 A (cont.)

MSRV pol cons ADN 41,42,43 Consensus	GCTCTGCTCA CAGGAGTTA GATACTTAGG GCTAAAATTA TCCAAAGGC GCTCTGCTCA CAGGAGTTA GATACTTAGG GCTAAAATTA TCCAAAGGC GCTCTGCTCA CAGGAGTTA GATACTTAGG GCTAAAATTA TCCAAAGGC	850 208 850
MSRV pol cons ADN 41,42,43 Consensus	CCAGGGCCCT CAGTGAGGA CGTATCCAGC CTATACTGG CTATACTCAT CCAGGGCCCT CAGTGAGGA CGTATCCAGC CTATACTGG CTATACTCAT CCAGGGCCCT CAGTGAGGA CGTATCCAGC CTATACTGG CTATACTCAT	900 258 900
MSRV pol cons ADN 41,42,43 Consensus	CCCCAAAACCC TAAAGCAACT AAAGAGGTTC CTGGCATAA CAGGTTTC CCCCAAAACCC TAAAGCAACT AAAGAGGTTC CTGGCATAA CAGGTTTC CCCCAAAACCC TAAAGCAACT AAAGAGGTTC CTGGCATAA CAGGTTTC	950 308 950
MSRV pol cons ADN 41,42,43 Consensus	CCAAAATACAG ATTCCCAGGT ACACGCCANT ACCCAGACCA TTAAATACAC CCAAAATACAG ATTCCCAGGT ACACGCCART ACCCAGACCA TTAAATACAC CCAAAATACAG ATTCCCAGGT ACACGCCMRNT ACCCAGACCA TTAAATACAC	1000 358 1000
MSRV pol cons ADN 41,42,43 Consensus	TAATTAGGAA AACTCAAAA GCCATTACCT ATTTAGTAAG ATGGACACT TAATTAGGAA AACTCAAAA GCCATTACCT ATTTAGTAAG ATGGACACT TAATTAGGAA AACTCAAAA GCCATTACCT ATTTAGTAAG ATGGACACT	1050 408 1050
MSRV pol cons ADN 41,42,43 Consensus	- CAGAAC TGGCTTTCGA GGCCCTAAAG AAGGCCCTAA CCCAAGCCCC GAACAGAAC TGGCTTTCGA GGCCCTAAAG - - - - GAACAGAAC TGGCTTTCGA GGCCCTAAAG AAGGCCCTAA CCCAAGCCCC	1097 438 1100
MSRV pol cons ADN 41,42,43 Consensus	AGTGTTCAGC TTGCCAACAG GGCAAGATT TTCTTTATAT GCCACAGAAA - - - - AGTGTTCAGC TTGCCAACAG GGCAAGATT TTCTTTATAT GCCACAGAAA	1147 438 1150
MSRV pol cons ADN 41,42,43 Consensus	AAACAGGAAT AGCTCTAGGA GTCCCTAACCG AGGTCTCAGG GATGAGCTTG - - - - AAACAGGAAT AGCTCTAGGA GTCCCTAACCG AGGTCTCAGG GATGAGCTTG	1197 438 1200
MSRV pol cons ADN 41,42,43 Consensus	CAACCCGTGG TATACCTGAG TAAGGAATT GATGTAGTGG CAAAGGGTTG - - - - CAACCCGTGG TATACCTGAG TAAGGAATT GATGTAGTGG CAAAGGGTTG	1247 438 1250
MSRV pol cons ADN 41,42,43 Consensus	GCCTCATTTGT TTATGGGTAA TGGGGCAGT AGCAGTCCTA GTATCTGAAG - - - - GCCTCATTTGT TTATGGGTAA TGGGGCAGT AGCAGTCCTA GTATCTGAAG	1297 438 1300
MSRV pol cons ADN 41,42,43 Consensus	CAGTAAAAAT AATACAGGGA AGAGATCTTA CTGTGTGGAC ATCTCATGAT - - - - CAGTAAAAAT AATACAGGGA AGAGATCTTA CTGTGTGGAC ATCTCATGAT	1347 438 1350
MSRV pol cons ADN 41,42,43 Consensus	GTGAACGGCA TACTCACTGC TAAAGGAGAC TTGTGGTTGT CAGACAACCA - - - - GTGAACGGCA TACTCACTGC TAAAGGAGAC TTGTGGTTGT CAGACAACCA	1397 438 1400
MSRV pol cons ADN 41,42,43 Consensus	TTTACTTAAT TATCAGGCTC TATTACTTGA AGAGCCAGTG CTGAGACTGC - - - - TTTACTTAAT TATCAGGCTC TATTACTTGA AGAGCCAGTG CTGAGACTGC	1447 438 1450
MSRV pol cons ADN 41,42,43 Consensus	GCACTTGTGC AACTCTTAAA CCCGCCACAT TTCTTCCAGA CAATGAAGAA - - - - GCACTTGTGC AACTCTTAAA CCCGCCACAT TTCTTCCAGA CAATGAAGAA	1497 438 1500
MSRV pol cons ADN 41,42,43 Consensus	AAGATAGAAC ATAACGTCA ACAAGTAATT GCTCAAACCT ATGCTGCTCG - - - - AAGATAGAAC ATAACGTCA ACAAGTAATT GCTCAAACCT ATGCTGCTCG	1547 438 1550
MSRV pol cons ADN 41,42,43 Consensus	AGGGGACCTT CTAGAGGTTC CCTTGACTGA TCCCGACCTC AACTGTATA - - - - AGGGGACCTT CTAGAGGTTC CCTTGACTGA TCCCGACCTC AACTGTATA	1597 438 1600

68/69

## FIG 52 B

Trans of MSRV pol cons prot 41,42,43	IMPESPTPLL GRDILAKAGA IIHLNIGKGI PICCPLLEEG INPEVWAIEG	50
Consensus	.....	50
Trans of MSRV pol cons prot 41,42,43	QYGQAKNARP VOVKLKDSAS FPYQRKYPLR PEALQGXQKI VKDLKAQGLV	100
Consensus	.....	100
Trans of MSRV pol cons prot 41,42,43	KPCSSPCNTP ILGVRKPNGQ WRLVODLRII NEAVFPLYP A VSSPYTLLSL	150
Consensus	.....	150
Trans of MSRV pol cons prot 41,42,43	IPEEEAEWFTV LDLKDAFFCI PVRPDSQFLF AFEDPLNPTS QLTWTIVLPQG	200
Consensus	.....	200
Trans of MSRV pol cons prot 41,42,43	FRDSPHLFGQ ALAQDLSQS YLDTLVLVYV DDILLI VARSE TLCHQATO[RE] [RE] DLSQS YLDTLVLVYV DDILLI VARSE TLCHQATO[RE] [RE] DLSQS YLDTLVLVYV DDILLI VARSE TLCHQATO[RE]	250
Consensus	.....	250
Trans of MSRV pol cons prot 41,42,43	[RE] [RE] TCGYK VSKPKAQLCS QEIHYLGLKL SKGTRNLSEE RIQPILGVPH [RE] [RE] TCGYK VSKPKAQLCS QEIHYLGLKL SKGTRNLSEE RIQPILGVPH [RE] [RE] TCGYK VSKPKAQLCS QEIHYLGLKL SKGTRNLSEE RIQPILGVPH	300
Consensus	.....	300
Trans of MSRV pol cons prot 41,42,43	PKTLKQLRGF LGITGFCKQ IPRYSHIARP LMTRIETQK ANIMLVRWTP PKTLKQLTAF LGITGFQIW IPRYSHIARP LMTRIETQK ANIMLVRWTP PKTLKQLTAF LGITGFQIW IPRYSHIARP LMTRIETQK ANIMLVRWTP	350
Consensus	.....	350
Trans of MSRV pol cons prot 41,42,43	TEVAFOAIKK ALTQAPVFSL PTGQDFSLYA TEKTGIALGV LTQVSGMSLQ	400
Consensus	.....	400
Trans of MSRV pol cons prot 41,42,43	PVVYLSKEID [RE] VAKGWPHEL WVMAAVAVLV SEAVKIIQGR DLTWWTSHDV [RE] FAE V [RE] [RE]	450
Consensus	.....	450
Trans of MSRV pol cons prot 41,42,43	NGILTAKGDL WLSDNHLLNY QAI LLEEPVL RLRTCATLKP ATFLPDNEEK [RE] AF QAI K [RE] [RE]	500
Consensus	.....	500
Trans of MSRV pol cons prot 41,42,43	IENHCQQVIA QTYAARGDLL EVPLTDPDLN LYTDGSSLAE KGLRKAGYAV	550
Consensus	.....	550
Trans of MSRV pol cons prot 41,42,43	ISDNGILESN RLTPGTS AHL AELIALTWAL ELGEGRKVNI YSDSKYAYLV	600
Consensus	.....	600
Trans of MSRV pol cons prot 41,42,43	LHAAHAAIWRE REFLTSEGTP INHQEAIRRL LLAVQKPKEV AVLHCQGHQE	650
Consensus	.....	650
Trans of MSRV pol cons prot 41,42,43	EEEREIEGNR QADIEAKKAA RQDSPLEMLI EGP	683
Consensus	.....	683

## FIG 53A

cons ADN 41,42,43	GACTTGAGCC AGTGTTCATA CCTGGACAMT CTTGTCATTC GGTACATGGG	50
cons ADN 1,5,8	GACTTGAGCC AGTGTTCATA CCTGGACAMT CTTGTCATTC GGTATTRGGGA	50
Consensus	GACTTGAGCC AGTGTTCATA CCTGGACAMT CTTGTCATTC GGTATTRGGGA	50
cons ADN 41,42,43	TGAAATTATTT TAGGCCACCC ATTCAAGAAC CTTGTCATTC CAAGCCACCC	100
cons ADN 1,5,8	TGAAATTATTT TAGGCCACCC ATTCAAGAAC CTTGTCATTC CAAGCCACCC	100
Consensus	TGAAATTATTT TAGGCCACCC ATTCAAGAAC CTTGTCATTC CAAGCCACCC	100
cons ADN 41,42,43	AAGCCTCTT AAATTTCCTT GCTACCTGTG GOTACAAGGT TICCAAACCA	150
cons ADN 1,5,8	AAGCCTCTT AAATTTCCTT GCTACCTGTG GOTACAAGGT TICCAAACCA	141
Consensus	AAGCCTCTT AAATTTCCTT GCTACCTGTG GOTACAAGGT TICCAAACCA	150
cons ADN 41,42,43	AGGGCTCAC TCTGCTCAC CAGGTAAA TACTTAGGGC TAAAATTATTC	200
cons ADN 1,5,8	AGGGCTCAC TCTGCTCAC CAGGTAAA TACTTAGGGC TAAAATTATTC	191
Consensus	AGGGCTCAC TCTGCTCAC CAGGTAAA TACTTAGGGC TAAAATTATTC	200
cons ADN 41,42,43	CAAAGGCCCC AGAACCTCTA GGAGGAACG TATCCAGGTT ATACTGGGTT	250
cons ADN 1,5,8	CAAAGGCCCC AGAACCTCTA GGAGGAACG TATCCAGGTT ATACTGGGTT	241
Consensus	CAAAGGCCCC AGAACCTCTA GGAGGAACG TATCCAGGTT ATACTGGGTT	250
cons ADN 41,42,43	ATCCCATCCC CAAACCTTA AAGCAACTAA CAGGTTCCCT TGGCATAACA	300
cons ADN 1,5,8	ATCCCATCCC CAAACCTTA AAGCAACTAA CAGGTTCCCT TGGCATAACA	291
Consensus	ATCCCATCCC CAAACCTTA AAGCAACTAA CAGGTTCCCT TGGCATAACA	300
cons ADN 41,42,43	GGTTTCTGCC PAATATGGAT TCCCGTAC AGCARRTAG CCAGGCCATT	350
cons ADN 1,5,8	GGTTTCTGCC PAATATGGAT TCCCGTAC AGCARRTAG CCAGGCCATT	341
Consensus	GGTTTCTGCC PAATATGGAT TCCCGTAC AGCARRTAG CCAGGCCATT	350
cons ADN 41,42,43	AAATACACCA ATTAAGGAAA CTCAAAAGC CATTACCCAT TTAGTAAGAT	400
cons ADN 1,5,8	AAATACACCA ATTAAGGAAA CTCAAAAGC CATTACCCAT TTAGTAAGAT	391
Consensus	AAATACACCA ATTAAGGAAA CTCAAAAGC CATTACCCAT TTAGTAAGAT	400
cons ADN 41,42,43	GGACACCTG ACCAGAAGTG GCTTTCCAGG CCCTAAAG	438
cons ADN 1,5,8	GGACACCTG ACCAGAAGTG GCTTTCCAGG CCCTAAAG	429
Consensus	GGACACCTG ACCAGAAGTG GCTTTCCAGG CCCTAAAG	438

## FIG 53B

cons prot 41,42,43	DLSQSSYLDI LVIHVNDLL IATHSETLH QATOALLNFL ATCGMKVSKP	50
cons prot 1,5,8	DLSQSSYLDI LVIHVNDLL IATHSETLH QATOALLNFL ATCGSK---Q	47
Consensus	DLSQSSYLDI LVIHVNDLL IATHSETLH QATOALLNFL ATCGK... .	50
cons prot 41,42,43	KAQLCSQQVK YLGLKLSKQT RNLREERIOP ILVYPHPKTL KQLTAFLGIT	100
cons prot 1,5,8	KAQLCSQQVK YLGLKLSKQT RNLREERIOP ILVYPHPKTL KQLRGFLGIT	97
Consensus	KAQLCSQQVK YLGLKLSKQT RNLREERIOP ILVYPHPKTL KQL... ELGIT	100
cons prot 41,42,43	GFCIWIPIRYSKIARPINIR IKETOKANTH VRWTPEPEEV AFQALK	146
cons prot 1,5,8	GFCIWIPIRYSKIARPINIR IKETOKANTH VRWTPEPEEV AFQALK	143
Consensus	GFCIWIPIRYSKIARPINIR IKETOKANTH VRWTPEPEEV AFQALK	146

# INTERNATIONAL SEARCH REPORT

International Application No

PCT/IB 97/01482

## A. CLASSIFICATION OF SUBJECT MATTER

IPC 6 C12N15/48	C12N5/08	C12N7/02	C07K14/15	C12N9/12
C12N9/22	C12Q1/70	C07K16/10	G01N33/569	A61K39/21
A61K39/42	A61K48/00			

According to International Patent Classification(IPC) or to both national classification and IPC

## B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

IPC 6 C12Q C07K C12N

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

## C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
A	EP 0 731 168 A (BIO MERIEUX) 11 September 1996 see the whole document ---	1-35
A	WO 95 21256 A (BIO MERIEUX ;PERRON HERVE (FR); MALLET FRANCOIS (FR); MANDRAND BER) 10 August 1995 see the whole document ---	1-35
A	WO 94 28138 A (UNIV LONDON ;GARSON JEREMY (GB); TUKE PHILIP (GB)) 8 December 1994 see the whole document ---	1-35

Further documents are listed in the continuation of box C.

Patent family members are listed in annex.

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Date of the actual completion of the international search

22 April 1998

Date of mailing of the international search report

08/05/1998

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International Application No

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## C.(Continuation) DOCUMENTS CONSIDERED TO BE RELEVANT

Category :	Citation of document, with indication where appropriate, of the relevant passages	Relevant to claim No.
A	PERRON H ET AL: "IN VITRO TRANSMISSION AND ANTIGENICITY OF A RETROVIRUS ISOLATED FROM A MULTIPLE SCLEROSIS PATIENT" RESEARCH IN VIROLOGY, vol. 143, no. 5, 1 January 1992, pages 337-350, XP000569296 see the whole document ---	1-35
P,X	PERRON ET AL.: "MOLECULAR IDENTIFICATION OF A NOVEL RETROVIRUS REPEATEDLY ISOLATED FROM PATIENTS WITH MULTIPLE SCLEROSIS" PNAS, vol. 94, July 1997, pages 7583-7588, XP002062853 see the whole document ---	1-35
P,A	WO 97 06260 A (BIO MERIEUX ;PERRON HERVE (FR); BESEME FREDERIC (FR); BEDIN FREDER) 20 February 1997 see the whole document -----	1-35

**INTERNATIONAL SEARCH REPORT**

Information on patent family members

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PCT/IB 97/01482

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